## Table 1: Genes in the Application

Function	UDP-N-ACETYLMURAMATEALANINE LIGASE (EC 6.3.2.8) UDP-N-ACETYLMURAMOYLALANINED-GLUTAMATE LIGASE (EC 6.3.2.9)	UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMATE2,6-DIAMINOPIMELATE	LIOANSE (EC 8.3.2.1.3) UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMYL-2,6-DIAMINOPIMELATED- ALANYL-D- ALANYL LIGASE	UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMYL-2,6-DIAMINOPIMELATED ALANYL-D- ALANYL LIGASE (EC 6.3.2.15)	UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMYL-2,6-DIAMINOPIMELATED- ALANYL-D-ALANYL LIGASE (FC 6.3.2.15)	UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMYL-2,6-DIAMINOPIMELATED- ALANYL-D-ALANYL LIGASE (FC 6.3.2.15)	UDP-N-ACETYLMURAMOYLALANYL-D-GĹUTAMATE2,6-DIAMINOPIMELATE LIGASE (EC 6.3.2.13)	FINE TANGLED PILI MAJOR SUBUNIT	NADPH DEHYDROGENASE 3 (EC 1.6.99.1)	NADPH-FLAVIN OXIDOREDUCTASE (EC 1.6.99)	SUPEROXIDE DISMUTASE [MN] (EC 1.15.1.1)	SUPEROXIDE DISMUTASE [MN] (EC 1.15.1.1)
NT Stop	115 4388	2539	18581	6920	7260	7723	8473	19557	602	10788	87008	87476
NT Start	1572 5803	3807	20110	7264	7694	8451	10035	19063	1573	9922	86877	87351
Contig.	GR00758 GR00758	GR00365	VV0017	GR00758	GR00758	GR00758	GR00758	00000	GR00259	VV0101	8600/\	VV0098
Identification Code	RXA02702 RXA02705	RXA01254	RXN02707	F RXA02707	F RXA02708	F RXA02709	RXA02710	RXN00531	RXA00944	RXS02560	RXS03119	RXS03120
Amino Acid	2 4	9	. σο	10	12	14	16	18	20	22	24	56
Nucleic Acid	) – 6	2	7	თ	=	13	15	17	19	21	23	25

### Cell wall biosynthesis

Function	N-ACETYLMURAMOYL-L-ALANINE AMIDASE (EC 3.5.1.28)	N-ACETYLMURAMOYL-L-ALANINE AMIDASE (EC 3.5.1.28)	UDP-N-ACETYLGLUCOSAMINE 1-CARBOXYVINYLTRANSFERASE (EC 2.5.1.7)	PHOSPHO-N-ACETYLMURAMOYL-PENTAPEPTIDE-TRANSFERASE (EC 2.7.8.13)	GLUTAMATE RACEMASE (EC 5.1.1.3)	D-ALANINED-ALANINE LIGASE (EC 6.3.2.4)	D-ALANINED-ALANINE LIGASE (EC 6.3.2.4)	UDP-N-ACETYLGLUCOSAMINEN-ACETYLMURAMYL-(PENTAPEPTIDE)	PYROPHOSPHORYL- UNDECAPRENOL N-ACETYLGLUCOSAMINE	TRANSFERASE (EC 2.4.1)	PENICILLIN-BINDING PROTEIN 2	
NT Stop	6271	3022	2962	5813	266	3381	806	1610		¢	10162	
NT Start	7458	2097	1709	6910	1845	4460	3	2698			12273	
Contig.	GR00417	GR00749	GR00021	GR00758	GR00703	VV0143	GR00292	GR00758			GR00758	
Identification Code	RXA01430	RXA02641	RXA00135	RXA02706	RXA02411	RXN01022		RXA02703	•		RXA02711	
Amino Acid	28	30	32	34	36	38	40	42		•3	44	t.
Nucleic Acid SEO ID NO	27	59	_31 _31	33	35	37	39	41			- 43	

Function Denicipal and Binding Obotein 6* DRECT IRSOR (D.A. ANY D.A. ANN)	CARBOXYPEPTIDASE) (EC 3.4.16.4)	PENICILLIN-BINDING PROTEIN 4	PENICILLIN-BINDING PROTEIN 1A	PENICILLIN-BINDING PROTEIN 1A	PENICILLIN-BINDING PROTEIN 3	PENICILLIN-BINDING PROTEIN 1A	PENICILLIN-BINDING PROTEIN 4 PRECURSOR (PBP-4) (D-ALANYL-D-ALANINE	CARBOXYPEPTIDASE) (EC 3.4.16.4) / D-ALANYL-D-ALANINE-ENDOPEPTIDASE (EC 3.4.99)	PENICILLIN-BINDING PROTEIN 5 PRECURSOR	(AL008883) penecillin binding protein (Mycobacterium tuberculosis)	perosamine synthetase	PENICILLIN-BINDING PROTEIN 1A	PENICILLIN-BINDING PROTEIN 1A	PENICILLIN-BINDING PROTEIN 1A	PENICILLIN-BINDING PROTEIN 5* PRECURSOR (D-ALANYL-D-ALANINE	CARBOXYPEPTIDASE) (EC 3.4.16.4)	PENICILLIN-BINDING PROTEIN 5* PRECURSOR (D-ALANYL-D-ALANINE	CARBOXYPEPTIDASE) (EC 3.4.16.4)	UDP-N-ACETYLGLUCOSAMINE 1-CARBOXYVINYLTRANSFERASE (EC 2.5.1.7)	UDP-N-acetylglucosamine-2-epimerase (EC 5.1.3.14) /N-acetylmannosamine kinase (EC 2.7.1.60)
NT Stop	7	4953	9561	4457	6315	1187	16650		5374	2675	20498	33419	33777	8970	121		121		16582	5317
NT Start	940	3928	10445	3525	7736	က	15370		3536	837	21652	31746	33457	9515	921		846		17895	4409
Contig.	GK 10003	GR00152	VV0054	GR00158	GR00516	GR00162	GR00424		VV0139	GR00449	GR00367	VV0079	620000	VV0054	VV0334		GR10005		6000/\	VV0119
Identification Code	KAAU2839	RXA00569	RXN03092	F RXA00594	RXA01828	RXA00612	RXA01510		RXN01608	F RXA01608	RXA01270	RXN00549	RXN00550	RXN03091	RXN03178		F RXA02859		RXN01267	RXN00045
Amino Acid SEQ ID NO	<del>Q</del>	48	20	52	54	26	58		09	62	64	99	99	20	72		74		76	78
Nucleic Acid SEQ ID NO	<b>c</b>	47	49	51	53	55	22		59	61	63	65	29	69	71		73		75	77

#### **Cell division**

Function	CELL DIVISIN PROTEIN FTSW	CELL DIVISIN PROTEIN FTSW	CELL DIVISION PROTEIN FTSZ	CELL DIVISION PROTEIN FTSX	CELL DIVISION ATP-BINDING PROTEIN FTSE	CELL DIVISION INHIBITOR	CELL DIVISION CONTROL PROTEIN 15 (EC 2.7.1)	CELL DIVISION PROTEIN FTSK	CELL DIVISION CONTROL PROTEIN 15 (EC 2.7.1)	CELL CYCLE PROTEIN MESJ	CELL DIVISION PROTEIN FTSH (EC 3.4.24)	CELL DIVISION PROTEIN FTSY	Hypothetical Cell Division Protein mraW	FTSQ	FTSQ	GLUCOSE INHIBITED DIVISION PROTEIN B
NT Stop	14355	2694	1404	646	1562	4847	5	1291	871	17596	20926	9069	13067	11080	2984	3403
NT Start	16043	4382	2729	1545	2248	6328	1588	2	2	16655	18368	4161	14077	11745	3460	2777
Contig.	VV0017	GR00758	GR00759	GR00002	GR00002	GR00022	GR00043	GR00233	GR00418	GR00424	GR00424	GR00630	GR00758	VV0017	GR00759	GR00417
Identification Code	RXN02704	F RXA02704	RXA02722	RXA00009	RXA00010	RXA00143	RXA00277	RXA00857	RXA01435	RXA01511	RXA01513	RXA02098	RXA02713	RXN02723	F RXA02723	RXA01426
Amino Acid SEQ ID NO	80	82	84	98	88	06	92	94	96	86	9	102	104	106	108	110
Nucleic Acid SEQ ID NO	6/	81	83	85	87	68	91	93	92	97	66	5	103	105	107	109

Function STAGE 0 SPORULATION PROTEIN J STAGE III SPORULATION PROTEIN E STAGE V SPORULATION PROTEIN E SOJ PROTEIN SOJ PROTEIN SOJ PROTEIN SOJ PROTEIN		티	ATP-DEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT CLPA	PEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT CLPA	ALP-DEPENDENT CLF PROTEASE ALP-BINDING SUBUNIT CLFA STOR DROTEIN	CLPB PROTEIN	/L-L-AMINO ACID AMIDOHYDROLASE (EC 3.5.1.14)	N-ACYL-L-AMINO ACID AMIDOHYDROLASE (EC 3.5.1.14)	/L-L-AMINO ACID AMIDOHYDROLASE (EC 3.5.1.14), hippurate hydrolase	[L42758) proteinase [Streptomyces lividans]	L42758) proteinase [Streptomyces lividans]	L42758) proteinase [Streptomyces lividans]	AMINOPERTIDAM AMINOPERTIDAM	OPEPTIDASE N (EC 3.4.11.2)	AMINOPEPTIDASE N (EC 3.4.11.2)	DPEPTIDASE N (EC 3.4.11.2)	ATP-DEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT CLPA	SEPENDENT OLY TROTEAGE ATT-BINDING SUBGINIT OLYA	DEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT CLPX	XAA-PRO AMINOPEPTIDASE (EC 3.4.11.9)	ZINC METALLOPROTEASE (EC 3.4.24)	METALLOPROTEASE (EC 3.4.24)	METALLOPROTEASE (EC 3.4.24)	PROLINE IMINOPEPTIDASE (EC 3.4.11.5)	INE IMINOPEPTIDASE	PROLYL ENDOPEPTIDASE (EC 3.4.21.26)	YL ENDOPEPTIDASE (EC 3.4.21.26)	IONINE AMINOPEPTIDASE (EC 3.4.11.18)	IONINE AMINOPEPTIDASE (EC 3.4.11.18)	PEPTIOYL-UIPEPTIDASE DCP (EC 3.4.15.5) PEPTIOYI - DIPEPTIDASE DCP (EC 3.4.15.5)	יסורטין איניין
STAGE (STAGE) STAGE (SOU PR(SOU PR(SU PR(SU)PR(SU PR(SU PRSUPRSU)PR(SU PRSUPRSU)PRSUPRSU)PR)))))))))))))))))))		Function	ATP-[	ATP-(	4 C	CLPB B	N-AC	N-AC	N-AC	(L427	(L427	(L427	Z	Ž Ž Ž	AMIN	AMIN	ATP-[	ATP	ATP-	XAA-F	ZINC	ZINC	ZINC	PROL	PR0	PROL	PROL	MET	METH	7. q	
NT Stop 5631 1344 7736 4432 4 14663 27685		NT Stop	43930	3196	4884 1391	3920	85382	2913	2877	6091	2660	4949	957	10728	1580	3152	49641	4401	1072	4603	11905	တ္တ	1652	9438	929	34158	20	34049	484	3885	3
NT Start 4495 4661 9058 3512 657 14043 28524		NT Start	41156	2216	1794	2205	85783	1729	1693	7596	1647	5194		13328	က	2289	47863	5678	2349	3803	9980	1640	1954	8158	က	32155	1738	33258	2	1222	,
Contig. GR00417 GR00516 GR00517 VV0229 GR00447 VV0054		Contig.	00000	GR00715	GK00/15	GR00464	8600/\	VV0043	GR10002	W0149	GR00275	GR00276	0 V V V V V V V V V V V V V V V V V V V	W0209	GR00289	GR00290	W0015	VV0210 VV0182	GR00310	VV0025	VV0127	GR00534	GR00534	080000	GR00125	6000	GR00368	VV0005	GR00178	VV0099	414000
Identification Code RXA01428 RXA01640 RXA01829 RXA01427 RXN02973 F RXA01603		Identification Code	RXN03028	F RXA02470	F KXAUZ4/1	F RXA01668	RXN02937	RXN03077	F RXA02855	RXN00982	F RXA00977	F RXA00982	F RXA01181	RXN01014	F RXA01014	F RXA01018	EXN01046	EXN01374	F RXA01120	RXN00397	RXN01868	F RXA01868	F RXA01869	RXN00499	F RXA00499	RXN01277	F RXA01277	RXN00675	F RXA00675	E RX A 00877	
Amino Acid SEQ ID NO 112 114 116 120 122	sis	Amino Acid	126	128	130	134	136	138	140	142	144	146 146	150	152	154	156	158	55	164	166	168	170	172	174	176	178	180	182	<u>7</u>	186 188	3
Nucleic Acid SEQ ID NO 111 113 115 117 119 121	Proteolysis	Nucleic Acid	125	127	129	133	135	137	139	141	143	145	149	151	153	155	157	151	163	165	167	169	171	173	175	177	179	181	183	185 187	2

Function PEPTIDYL-TRNA HYDROLASE (EC 3.1.1.29) Hypothetical Secretory Serine Protease (EC 3.4.21) PROTEASE II (EC 3.4.21.83) PTRB periplasmic protease PROTEASE II (EC 3.4.21.83) PTRB periplasmic protease PROTEIN P60 PRECURSOR HYDROGENASE 1 MATURATION PROTEASE (EC 3.4) GAMMA-GLUTAMYLTRANSPEPTIDASE (EC 2.3.2.2) GAMMA-GLUTAMYLTRANSPEPTIDASE (EC 2.3.2.2) GAMMA-GLUTAMYLTRANSPEPTIDASE (EC 2.3.2.2) GAMMA-GLUTAMYLTRANSPEPTIDASE (EC 2.3.2.2) GAMMA-GLUTAMYLTRANSPEPTIDASE PRECURSOR (EC 2.3.2.2) Bembrane Spanning Protease PEPTIDYL-TRNA HYDROLASE (EC 3.1.1.29) Serine protease
_,
NT Stop 4711 6 5071 4857 3735 6193 15368 40076 6109 507 3933 12074 1522 8139
NT Start 4172 689 5853 4075 5150 4778 14742 39393 4799 1 1 12751 2733 7528
Contig. VV0064 VV0200 VV0135 GR00163 VV0135 GR00163 VV0131 GR00801 GR00801 GR00801 GR00801 GR00801 VV0169
Identification Code RXN01226 RXN01963 RXN00621 F RXA00622 F RXA00622 F RXA00622 F RXA02820 F RXA02820 F RXA02820 F RXA02944 RXS01233 RXS01233 RXS01233
Amino Acid SEQ ID NO 190 192 194 196 198 200 202 204 208 210 212 214 216 218
Nucleic Acid SEQ ID NO 189 191 193 195 197 199 201 203 205 207 209 211 213

#### Enzymes in general

Function	BETA C-S LYASE (EC 3) PUTATIVE AMINOTRANSFERASE	Acetyltransferases	Acetyltransferases	ENTEROBACTIN SYNTHETASE COMPONENT F	Acetyltransferases (the isoleucine patch superfamily)	D-AMINO ACID DEHYDROGENASE LARGE SUBUNIT (EC 1.4.99.1)	D-AMINO ACID DEHYDROGENASE SMALL SUBUNIT (EC 1.4.99.1)	D-AMINO ACID DEHYDROGENASE SMALL SUBUNIT (EC 1.4.99.1)	2,3,4,5-tetrahydropyridine-2-carboxylate N-succinyltransferase (EC 2.3.1.117)	quinate dehydrogenase (pyrroloquinoline-quinone) (EC 1.1.99.25)	NITRILASE REGULATOR	Methyltransferase	QUINONE OXIDOREDUCTASE (EC 1.6.5.5)	3-KETOACYL-COA THIOLASE (EC 2.3.1.16)	UNDECAPRENYL-PHOSPHATE ALPHA-N-	ACETYLGLUCOSAMINYLTRANSFERASE (EC 2.4.1)	UNDECAPRENYL-PHOSPHATE GALACTOSEPHOSPHOTRANSFERASE (EC	2.7.8.6)	ACYLTRANSFERASE (EC 2.3.1)	DNA FOR L-PROLINE 3-HYDROXYLASE, COMPLETE CDS	ALCOHOL DEHYDROGENASE (EC 1.1.1.1)			
NT Stop	1478	1562	17387	3213	3213	5637	2	4	6046	4720	5053	1061		6460	4309	15596	10341	363		20990		15638	37450	44436
NT Start	2452	954	16827	7034	7034	3736	298	831	7548	4821	5952	2008		6930	3650	14586	9118	893		21430		16024	36800	43945
Contig.	GR00489	GR00650	GR00758	VV0008	GR00424	VV0321	GR00209	GR00210	GR00296	GR00296	GR00296	VV0160		W0196	VV0232	620070	VV0182	VV0268		6000//		VV0122	VV0139	6000/\
Identification Code	RXA01728	RXA02214	RXA02716	RXN01499	FRXA01499	RXN00787	F RXA00787	F RXA00791	RXA01057	RXA01055	RXA01056	RXN02021	RXS00949	RXS00004	RXS00166	RXS00288	RXS01114	RXS01205		RXS01269	-	RXS01421	RXS01491	RXS01572
Amino Acid	220	222	224	226	228	230	232	234	236	238	240	242	244	246	248	250	252	254		256		258	260	262
Nucleic Acid	219	221	223	225	227	229	231	233	235	237	239	241	243	245	247	249	251	253		255		257	259	261

			1.158)											
Function	ACETOIN(DIACETYL) REDUCTASE (EC 1.1.1.5)	(S,S)-butane-2,3-diol dehydrogenase (EC 1.1.1.76)	UDP-N-ACETYLENOLPYRUVOYLGLUCOSAMINE REDUCTASE (EC 1.1.1.158)	ALDEHYDE DEHYDROGENASE (EC 1.2.1.3)	ACYLTRANSFERASE	QUINONE OXIDOREDUCTASE (EC 1.6.5.5)	ALDEHYDE DEHYDROGENASE (EC 1.2.1.3)	ALDEHYDE DEHYDROGENASE (EC 1.2.1.3)	Oxidoreductase (EC 1.1.1)	METHYLTRANSFERASE (EC 2.1.1)	CAFFEOYL-COA O-METHYLTRANSFERASE (EC 2.1.1.104)	CAFFEOYL-COA O-METHYLTRANSFERASE (EC 2.1.1.104)	Protein involved in hydrolysis of epoxides	Metal-Dependent Hydrolase
NT Stop	8122	46248	3459	15815	6565	6733	437	10055				5057	26969	3715
NT Start	7370	47021	2359	17332	7668	5768	108	10678				4644	27517	4545
Contig.	VV0107	00000	70007	VV0057	8600//	VV0074	VV0034	VV0155				GR00549	VV0054	VV0105
Identification Code	RXS02453	RXS02474	RXS02485	RXS02539	RXS02578	RXS02741	RXS03061	RXS03150	RXS02554	RXS03058	RXS03218	F RXA01918	RXC00110	RXC01971
Amino Acid	264	266	268	270	272	274	276	278	280	282	284	286	288	290
Nucleic Acid	263	265	267	269	271	273	275	277	279	281	283	285	287	289

# Genes encoding enzymes for the metabolism of inorganic compounds

## Phosphate and Phosphonate metabolism

Function	PHNA PROTEIN	PHNB PROTEIN	PHNB PROTEIN	PHOH PROTEIN HOMOLOG	PHOH PROTEIN HOMOLOG	PHOH PROTEIN HOMOLOG	PHOSPHATE ACETYLTRANSFERASE (EC 2.3.1.8)	PHOSPHATE-BINDING PERIPLASMIC PROTEIN PRECURSOR	PHOSPHATE-BINDING PERIPLASMIC PROTEIN PRECURSOR	EXOPOLYPHOSPHATASE (EC 3.6.1.11)	ALKALINE PHOSPHATASE D PRECURSOR (EC 3.1.3.1)	INORGANIC PYROPHOSPHATASE (EC 3.6.1.1)	DEDA PROTEIN, similar to alkaline phosphatase	DEDA PROTEIN	DEDA PROTEIN - ALKALINE PHOSPHATASE LIKE PROTEIN	DEDA PROTEIN	CARBOXYVINYL-CARBOXYPHOSPHONATE PHOSPHORYLMUTASE (EC	2.7.8.23)	PHOSPHONATES TRANSPORT SYSTEM PERMEASE PROTEIN PHNE	PHOSPHONATES TRANSPORT ATP-BINDING PROTEIN PHNC	PHOSPHONATES TRANSPORT SYSTEM PERMEASE PROTEIN PHNE	alkaline phosphatase
NT Stop	1783	5962	4	11493	227	15341	2550	19250	8246	10985	10016	14696	10111	2774	1032	525	4260		6407	8055	7252	83037
NT Start	2124	6375	294	10120	1222	14325	3932	18126	9029	10059	8469	15169	9512	3355	286	79	5021		7252	8858	8055	84095
Contig.	GR00636	GR00012	GR00632	VV0142	GR00173	GR00242	GR00418	VV0103	GR00205	GR00720	GR00422	GR00424	GR00014	GR00162	VV0189	GR00602	GR00636		VV0106	W0106	VV0106	W0098
Identification Code	RXA02118	RXA00078	RXA02105	RXN00663	F RXA00663	RXA00888	RXA01437	RXN00778	F RXA00778	RXA02497	RXA01477	RXA01509	RXA00100	RXA00615	RXN00250	F RXA02010	RXA02120		RXS01000	RXS01002	RXS01003	RXS01902
Amino Acid SEQ ID NO	292	294	296	298	300	302	304	306	308	310	312	314	316	318	320	322	324		326	328	330	332
Nucleic Acid	291	293	295	297	299	301	303	305	307	309	311	313	315	317	319	321	323		325	327	329	331

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Attorney Docket No.: BGI-132CP

Function Function Reference Homeing Co.	FERRIC UPTAKE REGULATION PROTEIN	FERRIPYOCHELIN BINDING PROTEIN	FERRITIN	FERRITIN	IRON REPRESSOR	IRON(III) DICITRATE-BINDING PERIPLASMIC PROTEIN PRECURSOR	PERIPLASMIC-IRON-BINDING PROTEIN SHIB	FERRIC ANGUIBACTIN-BINDING PROTEIN PRECURSOR	FERROCHELATASE (EC 4.99.1.1)	FERROCHELATASE (EC 4.99.1.1)			
NT Stop	3867	7749	33793	935	827	1241	1757	3532	3795	2729	5402	2451	1332
NT Start	3436 3436	7192	33308	546	1486	2185	2692	2585	4586	1653	4389	1342	2018
O <sub>1</sub>	GR00011	_	VV0084	GR00586	GR00302	GR00358	GR00393	GR00451	GR00624	GR00078	GR00013	VV0167	
Identification Code	RXA00070	RXA01934	RXN01997	F RXA01997	RXA01082	RXA01236	RXA01354	RXA01620	RXA02052	RXA00372	RXA00088	RXS00156	RXS00624
Acid	336 336												
Nucleic Acid SEQ ID NO	335	337	339	341	343	345	347	349	351	353	355	357	359

# Modification and degradation of aromatic compounds

Function	ARYL-ALCOHOL DEHYDROGENASE (NADP+) (EC 1.1.1.91)	3-CARBOXY-CIS,CIS-MUCONATE CYCLOISOMERASE (EC 5.5.1.2)	3-CARBOXY-CIS, CIS-MUCONATE CYCLOISOMERASE HOMOLOG (EC 5.5.1.2)	3-CARBOXY-CIS, CIS-MUCONATE CYCLOISOMERASE HOMOLOG (EC 5.5.1.2)	4-CARBOXYMUCONOLACTONE DECARBOXYLASE (EC 4.1.1.44)	4-CARBOXYMUCONOLACTONE DECARBOXYLASE (EC 4.1.1.44)	MUCONATE CYCLOISOMERASE (EC 5.5.1.1)	MUCONATE CYCLOISOMERASE (EC 5.5.1.1)	MUCONOLACTONE ISOMERASE (EC 5.3.3.4)	4-HYDROXYBENZOATE OCTAPRENYLTRANSFERASE (EC 2.5.1)	4-HYDROXYBENZOATE OCTAPRENYLTRANSFERASE (EC 2.5.1)	4-HYDROXYBENZOATE OCTAPRENYLTRANSFERASE (EC 2.5.1)	BENZENE 1,2-DIOXYGENASE SYSTEM FERREDOXINNAD(+) REDUCTASE	COMPONENT (EC 1.18.1.3)	BIPHENYL-2,3-DIOL 1,2-DIOXYGENASE III (EC 1.13.11.39)	CAFFEOYL-COA O-METHYLTRANSFERASE (EC 2.1.1.104)	CATECHOL 1,2-DIOXYGENASE (EC 1.13.11.1)	CATECHOL 1,2-DIOXYGENASE (EC 1.13.11.1)	DIBENZOTHIOPHENE DESULFURIZATION ENZYME A	DIBENZOTHIOPHENE DESULFURIZATION ENZYME A	DIBENZOTHIOPHENE DESULFURIZATION ENZYME A	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE FORMING) 1 (EC 1.14.13.8)	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE FORMING) 1 (EC 1.14.13.8)	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE FORMING) 2 (EC 1.14.13.8)
NT Stop	1882	5314	14118	10	862	1876	2961	8025	2655	8737	449	419	9617		572	4585	8712	9	11407	804	971	6125	469	311
NT Start	938	4109	13120	651	1098	1556	4121	9038	2945	7742	۳	က	8385		15	5223	7858	665	12867	445	1909	5469	20	1720
Contig.	GR00003	GR00725	VV0128	GR00794	GR00307	GR00637	GR00421	GR00665	GR00421	GR00742	VV0362	GR00832	GR00424		GR00813	GR00626	VV0128	GR00168	VV0321	GR00212	GR00458	VV0057	GR00726	GR00629
Identification Code	RXA00024	RXA02526	RXN02813	F RXA02813	RXA01113	RXA02126	RXA01465	RXA02316	RXA01464	RXA02603	RXN02839	F RXA02839	RXA01502		RXA02828	RXA02064	RXN00639	F RXA00639	RXN01653	F RXA00797	F RXA01653	RXN02530	F RXA02530	RXA02083
Amino Acid	362	364	366	368	370	372	374	376	378	380	382	384	386		388	390	392	394	396	398	400	402	404	406
Nucleic Acid	361	363	365	367	369	371	373	375	377	379	381	383	385		387	389	391	393	395	397	388	401	403	405

Nicleic Acid	Amino Acid	Identification Code	Contin	Tets: LN	NT Ston	Finction
SEQ ID NO	SEQ ID NO		3		200	
407	408	RXA00892	GR00243	2188	1295	PARANITROBENZYL ESTERASE (EC 3.1.1)
409	410	RXA02092	GR00629	12153	10516	PARANITROBENZYL ESTERASE (EC 3.1.1)
411	412	RXN00658	VV0083	15705	16397	PHENOL 2-MONOOXYGENASE (EC 1.14.13.7)
413	414	F RXA00658	GR00170	321	4	PHENOL 2 MONOOXYGENASE (EC 1.14.13.7)
415	416	RXA01385	GR00406	5320	3440	PHENOL 2 MONOOXYGENASE (EC 1.14.13.7)
417	418	RXN01461	VV0128	12414	13025	PROTOCATECHUATE 3,4-DIOXYGENASE ALPHA CHAIN (EC 1.13.11.3)
419	420	F RXA01461	GR00421	463	2	PROTOCATECHUATE 3,4-DIOXYGENASE ALPHA CHAIN (EC 1.13.11.3)
421	422	RXA01462	GR00421	1167	478	PROTOCATECHUATE 3,4-DIOXYGENASE BETA CHAIN (EC 1.13.11.3)
423	424	RXN00641	W0128	7440	5950	TOLUATE 1,2-DIOXYGENASE ALPHA SUBUNIT (EC 1.14.12)
425	426	F RXA00640	GR00168	1083	1331	TOLUATE 1,2-DIOXYGENASE ALPHA SUBUNIT (EC 1.14.12)
427	428	F RXA00641	GR00168	1533	2573	TOLUATE 1,2-DIOXYGENASE ALPHA SUBUNIT (EC 1.14.12)
429	430	RXA00642	GR00168	2616	3107	TOLUATE 1,2-DIOXYGENASE BETA SUBUNIT (EC 1.14.12)
431	432	RXA00643	GR00168	3122	4657	TOLUATE 1,2-DIOXYGENASE ELECTRON TRANSFER COMPONENT
433	434	RXN01993	VV0182	16	1143	VANILLATE DEMETHYLASE (EC 1.14)
435	436	F RXA01993	GR00584	<b>-</b>	366	VANILLATE DEMETHYLASE (EC 1.14)
437	438	F RXA02012	GR00604	2	670	VANILLATE DEMETHYLASE (EC 1.14)
439	440	RXA01994	GR00584	373	1347	VANILLATE DEMETHYLASE OXIDOREDUCTASE (EC 1)
441	442	RXA02535	GR00726	629	7753	XYLENE MONOOXYGENASE ELECTRON TRANSFER COMPONENT
443	444	RXA00964	GR00269	1575	451	1-hydroxy-2-naphthoate 1,2-dioxygenase (EC 1.13.11.38)
445	446	RXN01466	VV0019	7050	6091	ARYLESTERASE (EC 3.1.1.2)
447	448	F RXA01466	GR00422	826	ည	ARYLESTERASE (EC 3.1.1.2)
449	450	RXN03036	VV0014	671	9	PROTOCATECHUATE 3,4-DIOXYGENASE BETA CHAIN (EC 1.13.11.3)
451	452	F RXA02895	GR10037	671	9	CHLOROCATECHOL 1,2-DIOXYGENASE (EC 1.13.11.1)
453	454	RXA02449	GR00710	1458	2360	hydroxyquinol 1,2-dioxygenase (EC 1.13.11.37)
455	456	RXN00178	VV0174	14670	15554	hydroxyquinol 1,2-dioxygenase (EC 1.13.11.37)
457	458	F RXA00178	GR00028	304	1188	HYDROXYQUINOL-1, 2-DIOXYGENASE
459	460	RXA02111	GR00632	4310	5593	QUINOLINATE SYNTHETASE A
461	462	RXA00644		GR0016	4657	CIS-1,2-DIHYDROXYCYCLOHEXA-3,5-DIENE-1-CARBOXYLATE
				80		DEHYDROGENASE (EC 1.3.1.55)
463	464	RXN00177	VV0174	13589	14656	MALEYLACETATE REDUCTASE (EC 1.3.1.32)
465	466	F RXA00177	GR00028	ဗ	290	MALEYLACETATE REDUCTASE (EC 1.3.1.32) metabolism of 2,4,5-
						trichlorophenoxyacetic acid
467	. 468	RXA02448	GR00710	340	1428	MALEYLACETATE REDUCTASE (EC 1.3.1.32)
469	470	RXA00048	GR00008	2185	527	3-(3-HYDROXYPHENYL) PROPIONATE HYDROXYLASE
471	472	RXA01126	GR00313	2	565	POSSIBLE 2-HYDROXYHEPTA-2,4-DIENE-1,7- DIOATE ISOMAERASE
473	474	RXA01117	GR00309	1713	973	SUCCINYL-COA:3-KETOACID-COENZYME A TRANSFERASE PRECURSOR (EC
						2.8.3.5)
475 477	476 478	RXA00772 RXA01288	GR00205 GR00372	2715 2018	1210 1644	SUCCINYL-COA:COENZYME A TRANSFERASE (EC 2.8.3) SUCCINYL-COA:COENZYME A TRANSFERASE (EC 2.8.3)
	) F		- : ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ;	?		

# TABLE 2: GENES IDENTIFIED FROM GENBANK

murC; ftsQ; ftsZ  murC; ftsQ; ftsZ  murC; ftsQ; ftsZ  murI Threonine dehydratase  dtsR1 dtsR2  murI D-glutamate racemase  ttr transketolase  gltB; gltD Glutamine 2-oxoglutarate aminotransferase  acn aconiase  rep, Replication protein; aminoglycoside  ach Argininosuccinate synthetase  glnA Glutamine synthetase  glnA Glutamine synthetase  argG Argininosuccinate synthetase  argG Argininosuccinate synthetase  argG Argininosuccinate synthetase  argG Argininosuccinate synthetase  argG Onnithine carbamolytransferase  argG Argininosuccinate synthetase  argG Onnithine carbamolytransferase  argG Onnithine carbamolytransferase  argG Onnithine carbamolytransferase	GenBank	Gene Name	Gene Function	Reference
Phosphoenol pyruvate carboxylase  Threonine dehydratase  Threonine dehydratase  Threonine dehydratase  Threonine dehydratase  Threonine dehydratase  Threonine dehydratase  Deglutamate racemase  Transketolase  Murl Transketolase  Harge and small subunits  aconitase  A acn aconitase  Herp Replication protein; aminoglycoside adenyltransferase  A acn adenyltransferase  Adenyltransferase  Blad Glutamine synthetase  Agent Glutamine synthetase  Blad Argininosuccinate synthetase  Argininosuccinate synthetase	Accession No.			
Threonine dehydratase	A09073	<b>8dd</b>	Phosphoenol pyruvate carboxylase	Bachmann, B. et al. "DNA fragment coding for phosphoenolpyruvat corboxylase, recombinant DNA carrying said fragment, strains carrying the recombinant DNA and method for producing L-aminino acids using said strains," Patent: EP 0358940-A 3 03/21/90
murC; ftsQ; ftsZ  dtsR1; dtsR2  murI D-glutamate racemase tkt transketolase gltB; gltD Glutamine 2-oxoglutarate aminotransferase large and small subunits acn aconitase rep, aad Replication protein; aminoglycoside adenyltransferase argC Replication protein; aminoglycoside adenyltransferase dehydrogenase glnA Glutamine synthetase argG Argininosuccinate synthetase argG Argininosuccinate synthetase argF Ornithine carbamolytransferase argF Ornithine carbamolytransferase argB Ornithine carbamolytransferase	A45579, A45581, A45583, A45585 A45587		Threonine dehydratase	Moeckel, B. et al. "Production of L-isoleucine by means of recombinant micro-organisms with deregulated threonine dehydratase," Patent: WO 9519442-A 5 07/20/95
dtsR  dtsR1; dtsR2  murI  tkt  transketolase  gltB; gltD  gltB; gltD  rep; aad  argc  argC  glnA  Glutamine synthetase  argC  Argininosuccinate synthetase  argC  Argininosuccinate synthetase  argC  Argininosuccinate synthetase  argC  Ormithine carbamolytransferase	AB003132	murC; ftsQ; ftsZ		Kobayashi, M. et al. "Cloning, sequencing, and characterization of the ftsZ gene from coryneform bacteria," <i>Biochem. Biophys. Res. Commun.</i> , 236(2):383-388 (1997)
dtsR1; dtsR2  murI  tkt  transketolase  gltB; gltD  gltB; gltD  acn  rep  rep  Replication protein  rep; aad  argC  glnA  Glutamine 2-oxoglutarate aminotransferase large and small subunits  aconitase  aconitase  Replication protein  Replication protein; aminoglycoside adenyltransferase  dehydrogenase  glnA  Glutamine synthetase hisF  cyclase  argG  Argininosuccinate synthetase  argF  Ornithine carbamolytransferase argF  Ornithine carbamolytransferase argF  Ornithine arbamolytransferase	AB015023	murC; ftsQ		Wachi, M. et al. "A murC gene from Coryneform bacteria," Appl. Microbiol. Biotechnol., 51(2):223-228 (1999)
dtsR1; dtsR2   D-glutamai     tkt   transketola     gltB; gltD   Glutamine     acn   aconitase     rep   Replication     rep; aad   Replication     rep; aad   Argininosu     glnA   Glutamine     hisF   cyclase     argG   Argininosu     argF   Ornithine o	AB018530	dtsR		Kimura, E. et al. "Molecular cloning of a novel gene, dtsR, which rescues the detergent sensitivity of a mutant derived from <i>Brevibacterium</i> lactofermentum," Biosci. Biotechnol. Biochem., 60(10):1565-1570 (1996)
tkt transketola gltB; gltD Glutamine gltB; gltD Glutamine large and s aconitase rep; aad Replication rep; aad Replication argC N-acetylglt Glutamine hisF cyclase argG Argininosu argF Ornithine of aroD 3-dehydroc	AB018531	dtsR1; dtsR2		
tkt         transketola           gltB; gltD         Glutamine           acn         aconitase           rep         Replication           rep; aad         Replication           argC         N-acetylglt           glnA         Glutamine           hisF         cyclase           argG         Argininosu           argF         Omithine on           aroD         3-dehydroc	AB020624	murl	D-glutamate racemase	
gltB; gltD         Glutamine           acn         aconitase           rep         Replication           rep; aad         Replication           argC         N-acetylglt           glnA         Glutamine           hisF         cyclase           argG         Argininosu           argF         Ornithine on           aroD         3-dehydroc	AB023377	tkt	transketolase	
acn         aconitase           rep         Replication           rep; aad         Replication           argC         N-acetylglu           glnA         dehydroger           hisF         cyclase           argG         Argininosu           argF         Ornithine c           aroD         3-dehydroq	AB024708	gltB; gltD	Glutamine 2-oxoglutarate aminotransferase large and small subunits	
rep         Replication           rep, aad         Replication           argC         N-acetylglu           glnA         Glutamine           hisF         cyclase           argG         Argininosu           argF         Ornithine c           aroD         3-dehydroq	AB025424	acn	aconitase	
rep; aad         Replication adenyltrans argC           glnA         Glutamine dehydrogen cyclase argG           argG         Argininosu argF           aroD         3-dehydroq	AB027714	rep	Replication protein	
argC glnA hisF argG argF aroD	AB027715	rep; aad	Replication protein; aminoglycoside adenyltransferase	
glnA hisF argG argF aroD	AF005242	argC	N-acetylglutamate-5-semialdehyde dehydrogenase	
hisF argG argF aroD	AF005635	glnA	Glutamine synthetase	
argG aroD	AF030405	hisF	cyclase	
argF aroD	AF030520	argG	Argininosuccinate synthetase	
aroD	AF031518	argF	Ornithine carbamolytransferase	
	AF036932	aroD	3-dehydroquinate dehydratase	

AF038651 dciAE; apt; rel Dipeptide-binding protein; adenine phosphoribosyltransferase; GTP pyrophosphokinase AF041436 argR Arginine repressor AF045998 impA Inositol monophosphate phosphatase AF048974 argC; argJ; argB; Arginine acetyltransferase; arginine acetyltransferase; arginine acetyltransferase; arginine acetyltransferase; arginine repressor; argininosuccinate lyase AF050109 inhA Eroylgutamate kinase; acetylornithine transminase; ornithine acetyltransferase; arginine repressor; argininosuccinate synthase; argininosuccinate synthase; argininosuccinate lyase AF050109 inhA Eroylgutamate synthase; argininosuccinate lyase AF051846 his A Phosphoribosyltransferase AF052652 metA Homoserine O-acetyltransferase AF053071 arob Dehydroquinate synthetase AF0653071 arob Dehydroquinate synthetase AF0653071 arob Dehydroquinate synthetase AF0660558 his Glutamine amidotransferase AF0614433 aroA Synthase AF114233 aroA Synthase L-aspartate-alpha-decarboxylase precurso		Gene Name	Gene Function	Reference
argR impA argC; argJ; argB; argC; argJ; argB; argC; argH inhA hisG hisA metA aroB hisH hisE aroA	+		Pyruvate carboxylase	
argR impA argC; argJ; argB; argC; argJ; argR; argC; argH inhA hisG hisA aroB aroB hisE aroA aroA panD		AE; apt; rel	Dipeptide-binding protein; adenine phosphoribosyltransferase; GTP pyrophosphokinase	Wehmeier, L. et al. "The role of the Corynebacterium glutamicum rel gene in (p)ppGpp metabolism," <i>Microbiology</i> , 144:1853-1862 (1998)
impA  argH, argC; argJ; argB; argC; argH inhA hisG hisA aroB hisH hisE aroA aroA panD		'R	Arginine repressor	
argC; argJ; argB; argC; argJ; argB; argG; argH inhA hisG hisA aroB aroB hisH hisE aroA aroA		Pd	Inositol monophosphate phosphatase	A PARTY OF THE PAR
argC; argJ; argB; argD; argF; argR; argG; argH inhA hisG hisA aroB hisH hisE aroA aroA		H	Argininosuccinate lyase	
argG; argH inhA hisG hisA aroB nisH hisE aroA aroA		;C; argJ; argB; D; argF; argR;	N-acetylglutamylphosphate reductase; ornithine acetyltransferase; N-	
inhA hisG hisA aroB hisH hisE aroA	arg	G; argH	acetylglutamate kinase; acetylornithine	
inhA hisG hisA aroB hisE aroA aroA panD			carbamovltransferase: arginine renressor:	
inhA hisG hisA aroB aroB hisH hisE aroA panD			argininosuccinate synthase;	
inhA hisG hisA aroB hisH hisE aroA panD			argininosuccinate lyase	
hisA metA aroB hisH aroA aroA		ıA	Enoyl-acyl carrier protein reductase	
hisA  aroB hisE aroA  aroA		G	ATP phosphoribosyltransferase	
aroB hisH hisE aroA panD		A	Phosphoribosylformimino-5-amino-1-	
metA aroB hisH aroA aroA panD			phosphoribosyl-4-imidazolecarboxamide	
aroA aroA panD			Isomerase	
aroB hisE aroA panD		t.A	Homoserine O-acetyltransferase	Park, S. et al. "Isolation and analysis of metA, a methionine biosynthetic gene encoding homoserine acetyltransferase in Corynebacterium glutamicum," Mol. Cells., 8(3):286-294 (1998)
hisE aroA panD		3B	Dehydroquinate synthetase	
hisE aroA panD		H	Glutamine amidotransferase	
aroA panD		ij	Phosphoribosyl-ATP- pyrophosphohydrolase	
panD		Ϋ́	5-enolpyruvylshikimate 3-phosphate synthase	
		On D	L-aspartate-alpha-decarboxylase precursor	Dusch, N. et al. "Expression of the Corynebacterium glutamicum panD gene encoding L-aspartate-alpha-decarboxylase leads to pantothenate overproduction in Escherichia coli," <i>Appl. Environ. Microbiol.</i> , 65(4)1530-1539 (1999)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
AF124518	aroD; aroE	3-dehydroquinase; shikimate dehydrogenase	
AF124600	aroC; aroK; aroB; pepQ	Chorismate synthase; shikimate kinase; 3-dehydroquinate synthase; putative cytoplasmic peptidase	
AF145897	inhA		
AF145898	inhA		
AJ001436	ectP	Transport of ectoine, glycine betaine, proline	Peter, H. et al. "Corynebacterium glutamicum is equipped with four secondary carriers for compatible solutes: Identification, sequencing, and characterization of the proline/ectoine uptake system, ProP, and the ectoine/proline/glycine betaine carrier, EctP." J. Bacteriol., 180(22):6005-6012 (1998)
AJ004934	барD	Tetrahydrodipicolinate succinylase (incomplete <sup>1</sup> )	Wehrmann, A. et al. "Different modes of diaminopimelate synthesis and their role in cell wall integrity: A study with Corynebacterium glutamicum," J. Bacteriol., 180(12):3159-3165 (1998)
AJ007732	ppc; secG; amt; ocd; soxA	Phosphoenolpyruvate-carboxylase; ?; high affinity ammonium uptake protein; putative ornithine-cyclodecarboxylase; sarcosine oxidase	
AJ010319	ftsY, glnB, glnD; srp; amtP	Involved in cell division; PII protein; uridylyltransferase (uridylyl-removing enzmye); signal recognition particle; low affinity ammonium uptake protein	Jakoby, M. et al. "Nitrogen regulation in Corynebacterium glutamicum; Isolation of genes involved in biochemical characterization of corresponding proteins," FEMS Microbiol., 173(2):303-310 (1999)
AJ132968	cat	Chloramphenicol aceteyl transferase	
AJ224946	овш	L-malate: quinone oxidoreductase	Molenaar, D. et al. "Biochemical and genetic characterization of the membrane-associated malate dehydrogenase (acceptor) from Corynebacterium glutamicum," Eur. J. Biochem., 254(2):395-403 (1998)
AJ238250	upu	NADH dehydrogenase	
AJ238703	porA	Porin	Lichtinger, T. et al. "Biochemical and biophysical characterization of the cell wall porin of Corynebacterium glutamicum: The channel is formed by a low molecular mass polypeptide," <i>Biochemistry</i> , 37(43):15024-15032 (1998)
D17429		Transposable element IS31831	Vertes, A.A. et al. "Isolation and characterization of IS31831, a transposable element from Corynebacterium glutamicum," Mol. Microbiol., 11(4):739-746 (1994)

GenBank <sup>TM</sup>	Gene Name	Gene Function	Reference
D84102	odhA	2-oxoglutarate dehydrogenase	Usuda, Y. et al. "Molecular cloning of the Corynebacterium glutamicum (Brevibacterium lactofermentum AJ12036) odhA gene encoding a novel type of 2-oxoglutarate dehydrogenase," <i>Microbiology</i> , 142:3347-3354 (1996)
E01358	hdh; hk	Homoserine dehydrogenase; homoserine kinase	Katsumata, R. et al. "Production of L-thereonine and L-isoleucine," Patent: JP 1987232392-A 1 10/12/87
E01359		Upstream of the start codon of homoserine kinase gene	Katsumata, R. et al. "Production of L-thereonine and L-isoleucine," Patent: JP 1987232392-A 2 10/12/87
E01375		Tryptophan operon	
E01376	ирГ.; ирЕ	Leader peptide; anthranilate synthase	Matsui, K. et al. "Tryptophan operon, peptide and protein coded thereby, utilization of tryptophan operon gene expression and production of tryptophan," Patent: JP 1987244382-A 1 10/24/87
E01377		Promoter and operator regions of tryptophan operon	Matsui, K. et al. "Tryptophan operon, peptide and protein coded thereby, utilization of tryptophan operon gene expression and production of tryptophan," Patent: JP 1987244382-A 1 10/24/87
E03937		Biotin-synthase	Hatakeyama, K. et al. "DNA fragment containing gene capable of coding biotin synthetase and its utilization," Patent: JP 1992278088-A 1 10/02/92
E04040		Diamino pelargonic acid aminotransferase	Kohama, K. et al. "Gene coding diaminopelargonic acid aminotransferase and desthiobiotin synthetase and its utilization," Patent: JP 1992330284-A 1 11/18/92
E04041		Desthiobiotinsynthetase	Kohama, K. et al. "Gene coding diaminopelargonic acid aminotransferase and desthiobiotin synthetase and its utilization," Patent: JP 1992330284-A 1 11/18/92
E04307		Flavum aspartase	Kurusu, Y. et al. "Gene DNA coding aspartase and utilization thereof," Patent: JP 1993030977-A 1 02/09/93
E04376		Isocitric acid lyase	Katsumata, R. et al. "Gene manifestation controlling DNA," Patent: JP 1993056782-A 3 03/09/93
E04377		Isocitric acid lyase N-terminal fragment	Katsumata, R. et al. "Gene manifestation controlling DNA," Patent: JP 1993056782-A 3 03/09/93
E04484		Prephenate dehydratase	Sotouchi, N. et al. "Production of L-phenylalanine by fermentation," Patent: JP 1993076352-A 2 03/30/93
E05108		Aspartokinase	Fugono, N. et al. "Gene DNA coding Aspartokinase and its use," Patent: JP 1993184366-A 1 07/27/93
E05112		Dihydro-dipichorinate synthetase	Hatakeyama, K. et al. "Gene DNA coding dihydrodipicolinic acid synthetase and its use," Patent: JP 1993184371-A 1 07/27/93

GenBank™	Gene Name	Gene Function	Reference
Accession No.			
E05776		Diaminopimelic acid dehydrogenase	Kobayashi, M. et al. "Gene DNA coding Diaminopimelic acid dehydrogenase and its use," Patent: JP 1993284970-A 1 11/02/93
E05779		Threonine synthase	Kohama, K. et al. "Gene DNA coding threonine synthase and its use," Patent: JP 1993284972-A 1 11/02/93
E06110		Prephenate dehydratase	Kikuchi, T. et al. "Production of L-phenylalanine by fermentation method," Patent: JP 1993344881-A 1 12/27/93
E06111		Mutated Prephenate dehydratase	Kikuchi, T. et al. "Production of L-phenylalanine by fermentation method," Patent: JP 1993344881-A 1 12/27/93
E06146		Acetohydroxy acid synthetase	Inui, M. et al. "Gene capable of coding Acetohydroxy acid synthetase and its use," Patent: JP 1993344893-A 1 12/27/93
E06825		Aspartokinase	Sugimoto, M. et al. "Mutant aspartokinase gene," patent: JP 1994062866-A 1 03/08/94
E06826		Mutated aspartokinase alpha subunit	Sugimoto, M. et al. "Mutant aspartokinase gene," patent: JP 1994062866-A 1 03/08/94
E06827		Mutated aspartokinase alpha subunit	Sugimoto, M. et al. "Mutant aspartokinase gene," patent: JP 1994062866-A 1 03/08/94
E07701	secY		Honno, N. et al. "Gene DNA participating in integration of membraneous protein to membrane," Patent: JP 1994169780-A 1 06/21/94
E08177		Aspartokinase	Sato, Y. et al. "Genetic DNA capable of coding Aspartokinase released from feedback inhibition and its utilization," Patent: JP 1994261766-A 1 09/20/94
E08178,		Feedback inhibition-released Aspartokinase	Sato, Y. et al. "Genetic DNA capable of coding Aspartokinase released from feadback inhihition and its utilization." Patent: ID 1004261766. A 1 00/2004
E08180, E08180, E08181, E08182			ICCUDACK IIIIIDIIIDII aliu its uliiizaliDii, Tatciit. J. 1774201700-A 1 07/20/74
E08232		Acetohydroxy-acid isomeroreductase	Inui, M. et al. "Gene DNA coding acetohydroxy acid isomeroreductase," Patent: JP 1994277067-A 1 10/04/94
E08234	дээs		Asai, Y. et al. "Gene DNA coding for translocation machinery of protein," Patent: JP 1994277073-A 1 10/04/94
E08643		FT aminotransferase and desthiobiotin synthetase promoter region	Hatakeyama, K. et al. "DNA fragment having promoter function in coryneform bacterium," Patent: JP 1995031476-A 1 02/03/95
E08646		Biotin synthetase	Hatakeyama, K. et al. "DNA fragment having promoter function in coryneform bacterium," Patent: JP 1995031476-A 1 02/03/95

GenBank <sup>TM</sup>	Gene Name	Gene Function	Reference
E08649		Aspartase	Kohama, K. et al "DNA fragment having promoter function in coryneform bacterium," Patent: JP 1995031478-A 1 02/03/95
E08900		Dihydrodipicolinate reductase	Madori, M. et al. "DNA fragment containing gene coding Dihydrodipicolinate acid reductase and utilization thereof," Patent: JP 1995075578-A 1 03/20/95
E08901		Diaminopimelic acid decarboxylase	Madori, M. et al. "DNA fragment containing gene coding Diaminopimelic acid decarboxylase and utilization thereof," Patent: JP 1995075579-A 1 03/20/95
E12594		Serine hydroxymethyltransferase	Hatakeyama, K. et al. "Production of L-trypophan," Patent: JP 1997028391-A 1 02/04/97
E12760, E12759, E12758		transposase	Moriya, M. et al. "Amplification of gene using artificial transposon," Patent: JP 1997070291-A 03/18/97
E12764		Arginyl-tRNA synthetase; diaminopimelic acid decarboxylase	Moriya, M. et al. "Amplification of gene using artificial transposon," Patent: JP 1997070291-A 03/18/97
E12767		Dihydrodipicolinic acid synthetase	Moriya, M. et al. "Amplification of gene using artificial transposon," Patent: JP 1997070291-A 03/18/97
E12770		aspartokinase	Moriya, M. et al. "Amplification of gene using artificial transposon," Patent: JP 1997070291-A 03/18/97
E12773		Dihydrodipicolinic acid reductase	Moriya, M. et al. "Amplification of gene using artificial transposon," Patent: JP 1997070291-A 03/18/97
E13655		Glucose-6-phosphate dehydrogenase	Hatakeyama, K. et al. "Glucose-6-phosphate dehydrogenase and DNA capable of coding the same," Patent: JP 1997224661-A 1 09/02/97
L01508	IIvA	Threonine dehydratase	Moeckel, B. et al. "Functional and structural analysis of the threonine dehydratase of Corynebacterium glutamicum," J. Bacteriol., 174:8065-8072 (1992)
L07603	EC 4.2.1.15	3-deoxy-D-arabinoheptulosonate-7- phosphate synthase	Chen, C. et al. "The cloning and nucleotide sequence of Corynebacterium glutamicum 3-deoxy-D-arabinoheptulosonate-7-phosphate synthase gene," <i>FEMS Microbiol. Lett.</i> , 107:223-230 (1993)
L09232	IIvB; iIvN; iIvC	Acetohydroxy acid synthase large subunit; Acetohydroxy acid synthase small subunit; Acetohydroxy acid isomeroreductase	Keilhauer, C. et al. "Isoleucine synthesis in Corynebacterium glutamicum: molecular analysis of the ilvB-ilvN-ilvC operon," J. Bacteriol., 175(17):5595-5603 (1993)

GenBank™ Accession No.	Gene Name	Gene Function	Reference
L18874	PtsM	Phosphoenolpyruvate sugar phosphotransferase	Fouet, A et al. "Bacillus subtilis sucrose-specific enzyme II of the phosphotransferase system: expression in Escherichia coli and homology to enzymes II from enteric bacteria," PNAS USA, 84(24):8773-8777 (1987); Lee, J.K. et al. "Nucleotide sequence of the gene encoding the Corynebacterium glutamicum mannose enzyme II and analyses of the deduced protein sequence," FEMS Microbiol. Lett., 119(1-2):137-145 (1994)
L27123	aceB	Malate synthase	Lee, H-S. et al. "Molecular characterization of aceB, a gene encoding malate synthase in Corynebacterium glutamicum," J. Microbiol. Biotechnol., 4(4):256-263 (1994)
L27126		Pyruvate kinase	Jetten, M. S. et al. "Structural and functional analysis of pyruvate kinase from Corynebacterium glutamicum," <i>Appl. Environ. Microbiol.</i> , 60(7):2501-2507 (1994)
L28760	aceA	Isocitrate lyase	
L35906	dtxr	Diphtheria toxin repressor	Oguiza, J.A. et al. "Molecular cloning, DNA sequence analysis, and characterization of the Corynebacterium diphtheriae dtxR from Brevibacterium lactofermentum," J. Bacteriol., 177(2):465-467 (1995)
M13774		Prephenate dehydratase	Follettie, M.T. et al. "Molecular cloning and nucleotide sequence of the Corynebacterium glutamicum pheA gene," J. Bacteriol., 167:695-702 (1986)
M16175	5S rRNA		Park, Y-H. et al. "Phylogenetic analysis of the coryneform bacteria by 56 rRNA sequences," J. Bacteriol, 169:1801-1806 (1987)
M16663	дфл	Anthranilate synthase, 5' end	Sano, K. et al. "Structure and function of the trp operon control regions of Brevibacterium lactofermentum, a glutamic-acid-producing bacterium," <i>Gene</i> , 52:191-200 (1987)
M16664	ирА	Tryptophan synthase, 3'end	Sano, K. et al. "Structure and function of the trp operon control regions of Brevibacterium lactofermentum, a glutamic-acid-producing bacterium," <i>Gene</i> , 52:191-200 (1987)
M25819		Phosphoenolpyruvate carboxylase	O'Regan, M. et al. "Cloning and nucleotide sequence of the Phosphoenolpyruvate carboxylase-coding gene of Corynebacterium glutamicum ATCC13032," Gene, 77(2):237-251 (1989)
M85106	,	23S rRNA gene insertion sequence	Roller, C. et al. "Gram-positive bacteria with a high DNA G+C content are characterized by a common insertion within their 23S rRNA genes," J. Gen. Microbiol., 138:1167-1175 (1992)

GenBank™ Accession No.	Gene Name	Gene Function	Reference
M85107, M85108		23S rRNA gene insertion sequence	Roller, C. et al. "Gram-positive bacteria with a high DNA G+C content are characterized by a common insertion within their 23S rRNA genes," J. Gen. Microbiol., 138:1167-1175 (1992)
M89931	aecD; brnQ; yhbw	Beta C-S lyase; branched-chain amino acid uptake carrier; hypothetical protein yhbw	Rossol, I. et al. "The Corynebacterium glutamicum aecD gene encodes a C-S lyase with alpha, beta-elimination activity that degrades aminoethylcysteine," <i>J. Bacteriol.</i> , 174(9):2968-2977 (1992); Tauch, A. et al. "Isoleucine uptake in Corynebacterium glutamicum ATCC 13032 is directed by the brnQ gene product," <i>Arch. Microbiol.</i> , 169(4):303-312 (1998)
S59299	пр	Leader gene (promoter)	Herry, D.M. et al. "Cloning of the trp gene cluster from a tryptophan-hyperproducing strain of Corynebacterium glutamicum: identification of a mutation in the trp leader sequence," <i>Appl. Environ. Microbiol.</i> , 59(3):791-799 (1993)
U11545	Стр	Anthranilate phosphoribosyltransferase	O'Gara, J.P. and Dunican, L.K. (1994) Complete nucleotide sequence of the Corynebacterium glutamicum ATCC 21850 tpD gene." Thesis, Microbiology Department, University College Galway, Ireland.
U13922	cgIIM; cgIIR; clgIIR	Putative type II 5-cytosoine methyltransferase; putative type II restriction endonuclease; putative type I or type III restriction endonuclease	Schafer, A. et al. "Cloning and characterization of a DNA region encoding a stress-sensitive restriction system from Corynebacterium glutamicum ATCC 13032 and analysis of its role in intergeneric conjugation with Escherichia coli," J. Bacteriol., 176(23):7309-7319 (1994); Schafer, A. et al. "The Corynebacterium glutamicum cglIM gene encoding a 5-cytosine in an McrBC-deficient Escherichia coli strain," Gene, 203(2):95-101 (1997)
U14965	recA		
U31224	xdd		Ankri, S. et al. "Mutations in the Corynebacterium glutamicumproline biosynthetic pathway: A natural bypass of the proA step," J. Bacteriol., 178(15):4412-4419 (1996)
U31225	proC	L-proline: NADP+ 5-oxidoreductase	Ankri, S. et al. "Mutations in the Corynebacterium glutamicumproline biosynthetic pathway: A natural bypass of the proA step," J. Bacteriol., 178(15):4412-4419 (1996)
U31230	obg; proB; unkdh	?;gamma glutamyl kinase;similar to D- isomer specific 2-hydroxyacid dehydrogenases	Ankri, S. et al. "Mutations in the Corynebacterium glutamicumproline biosynthetic pathway: A natural bypass of the proA step," J. Bacteriol., 178(15):4412-4419 (1996)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
U31281	bioB	Biotin synthase	Serebriiskii, I.G., "Two new members of the bio B superfamily: Cloning, sequencing and expression of bio B genes of Methylobacillus flagellatum and Corynebacterium glutamicum," <i>Gene</i> , 175:15-22 (1996)
U35023	thtR; accBC	Thiosulfate sulfurtransferase; acyl CoA carboxylase	Jager, W. et al. "A Corynebacterium glutamicum gene encoding a two-domain protein similar to biotin carboxylases and biotin-carboxyl-carrier proteins," <i>Arch. Microbiol.</i> , 166(2);76-82 (1996)
U43535	ст	Multidrug resistance protein	Jager, W. et al. "A Corynebacterium glutamicum gene conferring multidrug resistance in the heterologous host Escherichia coli," J. Bacteriol., 179(7):2449-2451 (1997)
U43536	clpB	Heat shock ATP-binding protein	
U53587	aphA-3	3.5"-aminoglycoside phosphotransferase	
U89648		Corynebacterium glutamicum unidentified sequence involved in histidine biosynthesis, partial sequence	
X04960	սթA; սթB; սթС; սթD; սթE; սթG; սթL	Tryptophan operon	Matsui, K. et al. "Complete nucleotide and deduced amino acid sequences of the Brevibacterium lactofermentum tryptophan operon," <i>Nucleic Acids Res.</i> , 14(24):10113-10114 (1986)
X07563	lys A	DAP decarboxylase (meso-diaminopimelate decarboxylase, EC 4.1.1.20)	Yeh, P. et al. "Nucleic sequence of the lysA gene of Corynebacterium glutamicum and possible mechanisms for modulation of its expression," Mol. Gen. Genet., 212(1):112-119 (1988)
X14234	EC 4.1.1.31	Phosphoenolpyruvate carboxylase	Eikmanns, B.J. et al. "The Phosphoenolpyruvate carboxylase gene of Corynebacterium glutamicum: Molecular cloning, nucleotide sequence, and expression," <i>Mol. Gen. Genet.</i> , 218(2):330-339 (1989); Lepiniec, L. et al. "Sorghum Phosphoenolpyruvate carboxylase gene family: structure, function and molecular evolution," <i>Plant. Mol. Biol.</i> , 21 (3):487-502 (1993)
X17313	fda	Fructose-bisphosphate aldolase	Von der Osten, C.H. et al. "Molecular cloning, nucleotide sequence and fine- structural analysis of the Corynebacterium glutamicum fda gene: structural comparison of C. glutamicum fructose-1, 6-biphosphate aldolase to class I and class II aldolases," Mol. Microbiol.,
X53993	dapA	L-2, 3-dihydrodipicolinate synthetase (EC 4.2.1.52)	Bonnassie, S. et al. "Nucleic sequence of the dapA gene from Corynebacterium glutamicum," Nucleic Acids Res., 18(21):6421 (1990)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
X54223		AttB-related site	Cianciotto, N. et al. "DNA sequence homology between att B-related sites of Corynebacterium diphtheriae, Corynebacterium ulcerans, Corynebacterium glutamicum, and the attP site of lambdacorynephage," FEMS. Microbiol, Lett., 66:299-302 (1990)
X54740	argS; lysA	Arginyl-tRNA synthetase; Diaminopimelate decarboxylase	Marcel, T. et al. "Nucleotide sequence and organization of the upstream region of the Corynebacterium glutamicum lysA gene," Mol. Microbiol., 4(11):1819-1830 (1990)
X55994	upL; trpE	Putative leader peptide; anthranilate synthase component 1	Heery, D.M. et al. "Nucleotide sequence of the Corynebacterium glutamicum trpE gene," <i>Nucleic Acids Res.</i> , 18(23):7138 (1990)
X56037	thrC	Threonine synthase	Han, K.S. et al. "The molecular structure of the Corynebacterium glutamicum threonine synthase gene," Mol. Microbiol., 4(10):1693-1702 (1990)
X56075	attB-related site	Attachment site	Cianciotto, N. et al. "DNA sequence homology between att B-related sites of Corynebacterium diphtheriae, Corynebacterium ulcerans, Corynebacterium glutamicum, and the attP site of lambdacorynephage," FEMS. Microbiol, Lett., 66:299-302 (1990)
X57226	lysC-alpha; lysC-beta; asd	Aspartokinase-alpha subunit; Aspartokinase-beta subunit; aspartate beta semialdehyde dehydrogenase	Kalinowski, J. et al. "Genetic and biochemical analysis of the Aspartokinase from Corynebacterium glutamicum," <i>Mol. Microbiol.</i> , 5(5):1197-1204 (1991); Kalinowski, J. et al. "Aspartokinase genes lysC alpha and lysC beta overlap and are adjacent to the aspertate beta-semialdehyde dehydrogenase gene asd in Corynebacterium glutamicum," <i>Mol. Gen. Genet.</i> , 224(3):317-324 (1990)
X59403	gap;pgk; tpi	Glyceraldehyde-3-phosphate; phosphoglycerate kinase; triosephosphate isomerase	Eikmanns, B.J. "Identification, sequence analysis, and expression of a Corynebacterium glutamicum gene cluster encoding the three glycolytic enzymes glyceraldehyde-3-phosphate dehydrogenase, 3-phosphoglycerate kinase, and triosephosphate isomeras," J. Bacteriol., 174(19):6076-6086 (1992)
X59404	qpg	Glutamate dehydrogenase	Bormann, E.R. et al. "Molecular analysis of the Corynebacterium glutamicum gdh gene encoding glutamate dehydrogenase," <i>Mol. Microbiol.</i> , 6(3):317-326 (1992)
X60312	lysI	L-lysine permease	Seep-Feldhaus, A.H. et al. "Molecular analysis of the Corynebacterium glutamicum lysl gene involved in lysine uptake," <i>Mol. Microbiol.</i> , 5(12):2995-3005 (1991)

GenBank <sup>TM</sup>	Gene Name	Gene Function	Reference
X66078	cop1	Ps1 protein	Joliff, G. et al. "Cloning and nucleotide sequence of the csp1 gene encoding PS1, one of the two major secreted proteins of Corynebacterium glutamicum: The deduced N-terminal region of PS1 is similar to the Mycobacterium antigen 85 complex," Mol. Microbiol., 6(16):2349-2362 (1992)
X66112	glt	Citrate synthase	Eikmanns, B.J. et al. "Cloning sequence, expression and transcriptional analysis of the Corynebacterium glutamicum gltA gene encoding citrate synthase," <i>Microbiol.</i> , 140:1817-1828 (1994)
X67737	dapB	Dihydrodipicolinate reductase	
X69103	csp2	Surface layer protein PS2	Peyret, J.L. et al. "Characterization of the cspB gene encoding PS2, an ordered surface-layer protein in Corynebacterium glutamicum," <i>Mol. Microbiol.</i> , 9(1):97-109 (1993)
X69104		IS3 related insertion element	Bonamy, C. et al. "Identification of IS1206, a Corynebacterium glutamicum IS3-related insertion sequence and phylogenetic analysis," <i>Mol. Microbiol.</i> , 14(3):571-581 (1994)
X70959	leuA	Isopropylmalate synthase	Patek, M. et al. "Leucine synthesis in Corynebacterium glutamicum: enzyme activities, structure of leuA, and effect of leuA inactivation on lysine synthesis," Appl. Environ. Microbiol., 60(1):133-140 (1994)
X71489	icd	Isocitrate dehydrogenase (NADP+)	Eikmanns, B.J. et al. "Cloning sequence analysis, expression, and inactivation of the Corynebacterium glutamicum icd gene encoding isocitrate dehydrogenase and biochemical characterization of the enzyme," J. Bacteriol., 177(3):774-782 (1995)
X72855	GDHA	Glutamate dehydrogenase (NADP+)	
X75083, X70584	mtrA	5-methyltryptophan resistance	Heery, D.M. et al. "A sequence from a tryptophan-hyperproducing strain of Corynebacterium glutamicum encoding resistance to 5-methyltryptophan," <i>Biochem. Biophys. Res. Commun.</i> , 201(3):1255-1262 (1994)
X75085	recA		Fitzpatrick, R. et al. "Construction and characterization of recA mutant strains of Corynebacterium glutamicum and Brevibacterium lactofermentum," Appl. Microbiol. Biotechnol., 42(4):575-580 (1994)
X75504	aceA; thiX	Partial Isocitrate lyase; ?	Reinscheid, D.J. et al. "Characterization of the isocitrate lyase gene from Corynebacterium glutamicum and biochemical analysis of the enzyme," J. Bacteriol., 176(12):3474-3483 (1994)
X76875		ATPase beta-subunit	Ludwig, W. et al. "Phylogenetic relationships of bacteria based on comparative sequence analysis of elongation factor Tu and ATP-synthase beta-subunit genes," Antonie Van Leeuwenhoek, 64:285-305 (1993)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
X77034	tuf	Elongation factor Tu	Ludwig, W. et al. "Phylogenetic relationships of bacteria based on comparative sequence analysis of elongation factor Tu and ATP-synthase beta-subunit genes," Antonie Van Leeuwenhoek, 64:285-305 (1993)
X77384	recA		Billman-Jacobe, H. "Nucleotide sequence of a recA gene from Corynebacterium glutamicum," DNA Seq., 4(6):403-404 (1994)
X78491	aceB	Malate synthase	Reinscheid, D.J. et al. "Malate synthase from Corynebacterium glutamicum pta-ack operon encoding phosphotransacetylase: sequence analysis," <i>Microbiology</i> , 140:3099-3108 (1994)
X80629	16S rDNA	16S ribosomal RNA	Rainey, F.A. et al. "Phylogenetic analysis of the genera Rhodococcus and Norcardia and evidence for the evolutionary origin of the genus Norcardia from within the radiation of Rhodococcus species," <i>Microbiol.</i> , 141:523-528 (1995)
X81191	gluA; gluB; gluC; gluD	Glutamate uptake system	Kronemeyer, W. et al. "Structure of the gluABCD cluster encoding the glutamate uptake system of Corynebacterium glutamicum," J. Bacteriol., 177(5):1152-1158 (1995)
X81379	dapE	Succinyldiaminopimelate desuccinylase	Wehrmann, A. et al. "Analysis of different DNA fragments of Corynebacterium glutamicum complementing dapE of Escherichia coli," <i>Microbiology</i> , 40:3349-56 (1994)
X82061	16S rDNA	16S ribosomal RNA	Ruimy, R. et al. 'Phylogeny of the genus Corynebacterium deduced from analyses of small-subunit ribosomal DNA sequences," <i>Int. J. Syst. Bacteriol.</i> , 45(4):740-746 (1995)
X82928	asd; lysC	Aspartate-semialdehyde dehydrogenase; ?	Serebrijski, I. et al. "Multicopy suppression by asd gene and osmotic stress-dependent complementation by heterologous proA in proA mutants," J. Bacteriol., 177(24):7255-7260 (1995)
X82929	proA	Gamma-glutamyl phosphate reductase	Screbrijski, I. et al. "Multicopy suppression by asd gene and osmotic stress-dependent complementation by heterologous proA in proA mutants," J. Bacteriol., 177(24):7255-7260 (1995)
X84257	16S rDNA	16S ribosomal RNA	Pascual, C. et al. "Phylogenetic analysis of the genus Corynebacterium based on 16S rRNA gene sequences," <i>Int. J. Syst. Bacteriol.</i> , 45(4):724-728 (1995)
X85965	aroP; dapE	Aromatic amino acid permease; ?	Wehrmann, A. et al. "Functional analysis of sequences adjacent to dapE of Corynebacterium glutamicumproline reveals the presence of aroP, which encodes the aromatic amino acid transporter," J. Bacteriol., 177(20):5991-5993 (1995)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
X86157	argB; argC; argD; argF; argJ	Acetylglutamate kinase; N-acetyl-gammaglutamyl-phosphate reductase; acetylornithine aminotransferase; ornithine carbamoyltransferase; glutamate N-acetyltransferase	Sakanyan, V. et al. "Genes and enzymes of the acetyl cycle of arginine biosynthesis in Corynebacterium glutamicum: enzyme evolution in the early steps of the arginine pathway," <i>Microbiology</i> , 142:99-108 (1996)
X89084	pta; ackA	Phosphate acetyltransferase; acetate kinase	Reinscheid, D.J. et al. "Cloning, sequence analysis, expression and inactivation of the Corynebacterium glutamicum pta-ack operon encoding phosphotransacetylase and acetate kinase," <i>Microbiology</i> , 145:503-513 (1999)
X89850	attB	Attachment site	Le Marrec, C. et al. "Genetic characterization of site-specific integration functions of phi AAU2 infecting "Arthrobacter aureus C70," J. Bacteriol., 178(7):1996-2004 (1996)
X90356		Promoter fragment F1	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90357		Promoter fragment F2	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90358		Promoter fragment F10	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90359		Promoter fragment F13	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90360		Promoter fragment F22	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90361		Promoter fragment F34	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90362		Promoter fragment F37	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
X90363		Promoter fragment F45	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90364		Promoter fragment F64	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90365		Promoter fragment F75	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90366		Promoter fragment PF101	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90367		Promoter fragment PF104	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X90368		Promoter fragment PF109	Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning, molecular analysis and search for a consensus motif," <i>Microbiology</i> , 142:1297-1309 (1996)
X93513	amt	Ammonium transport system	Siewe, R.M. et al. "Functional and genetic characterization of the (methyl) ammonium uptake carrier of Corynebacterium glutamicum," J. Biol. Chem., 271(10):5398-5403 (1996)
X93514	betP	Glycine betaine transport system	Peter, H. et al. "Isolation, characterization, and expression of the Corynebacterium glutamicum betP gene, encoding the transport system for the compatible solute glycine betaine," J. Bacteriol., 178(17):5229-5234 (1996)
X95649	orf4		Patek, M. et al. "Identification and transcriptional analysis of the dapB-ORF2-dapA-ORF4 operon of Corynebacterium glutamicum, encoding two enzymes involved in L-lysine synthesis," <i>Biotechnol. Lett.</i> , 19:1113-1117 (1997)
X96471	lysE; lysG	Lysine exporter protein; Lysine export regulator protein	Vrljic, M. et al. "A new type of transporter with a new type of cellular function: L-lysine export from Corynebacterium glutamicum," Mol. Microbiol., 22(5):815-826 (1996)

GenBank <sup>TM</sup> Accession No.	Gene Name	Gene Function	Reference
X96580	panB; panC; xylB	3-methyl-2-oxobutanoate hydroxymethyltransferase; pantoate-beta-alanine ligase; xylulokinase	Sahm, H. et al. "D-pantothenate synthesis in Corynebacterium glutamicum and use of panBC and genes encoding L-valine synthesis for D-pantothenate overproduction," <i>Appl. Environ. Microbiol.</i> , 65(5):1973-1979 (1999)
X96962		Insertion sequence IS1207 and transposase	
X99289		Elongation factor P	Ramos, A. et al. "Cloning, sequencing and expression of the gene encoding elongation factor P in the amino-acid producer Brevibacterium lactofermentum (Corynebacterium glutamicum ATCC 13869)," Gene, 198:217-222 (1997)
Y00140	thrB	Homoserine kinase	Mateos, L.M. et al. "Nucleotide sequence of the homoserine kinase (thrB) gene of the Brevibacterium lactofermentum," <i>Nucleic Acids Res.</i> , 15(9):3922 (1987)
Y00151	qpp	Meso-diaminopimelate D-dehydrogenase (EC 1.4.1.16)	Ishino, S. et al. "Nucleotide sequence of the meso-diaminopimelate D-dehydrogenase gene from Corynebacterium glutamicum," <i>Nucleic Acids Res.</i> , 15(9):3917 (1987)
Y00476	thrA	Homoserine dehydrogenase	Mateos, L.M. et al. "Nucleotide sequence of the homoserine dehydrogenase (thrA) gene of the Brevibacterium lactofermentum," <i>Nucleic Acids Res.</i> , 15(24):10598 (1987)
Y00546	hom; thrB	Homoserine dehydrogenase; homoserine kinase	Peoples, O.P. et al. "Nucleotide sequence and fine structural analysis of the Corynebacterium glutamicum hom-thrB operon," <i>Mol. Microbiol.</i> , 2(1):63-72 (1988)
Y08964	murC; ftsQ/divD; ftsZ	UPD-N-acetylmuramate-alanine ligase; division initiation protein or cell division protein; cell division protein	Honrubia, M.P. et al. "Identification, characterization, and chromosomal organization of the ftsZ gene from Brevibacterium lactofermentum," Mol. Gen. Genet., 259(1):97-104 (1998)
Y09163	putP	High affinity proline transport system	Peter, H. et al. "Isolation of the putP gene of Corynebacterium glutamicumproline and characterization of a low-affinity uptake system for compatible solutes," <i>Arch. Microbiol.</i> , 168(2):143-151 (1997)
Y09548	pyc	Pyruvate carboxylase	Peters-Wendisch, P.G. et al. "Pyruvate carboxylase from Corynebacterium glutamicum: characterization, expression and inactivation of the pyc gene," <i>Microbiology</i> , 144:915-927 (1998)
X09578	leuB	3-isopropylmalate dehydrogenase	Patek, M. et al. "Analysis of the leuB gene from Corynebacterium glutamicum," Appl. Microbiol. Biotechnol., 50(1):42-47 (1998)
Y12472		Attachment site bacteriophage Phi-16	Moreau, S. et al. "Site-specific integration of corynephage Phi-16: The construction of an integration vector," <i>Microbiol.</i> , 145:539-548 (1999)

GenBank <sup>TM</sup>	Gene Name	Gene Function	Reference
Accession No.	proP	Proline/ectoine uptake system protein	Peter, H. et al. "Corynebacterium glutamicum is equipped with four secondary carriers for compatible solutes: Identification, sequencing, and characterization of the proline/ertoine untake system. Prop. and the ectoine/proline/elvoine.
			betaine carrier, EctP., J. Bacteriol., 180(22):6005-6012 (1998)
Y13221	glnA	Glutamine synthetase I	Jakoby, M. et al. "Isolation of Corynebacterium glutamicum glnA gene encoding glutamine synthetase I," FEMS Microbiol. Lett., 154(1):81-88 (1997)
Y16642	lpd	Dihydrolipoamide dehydrogenase	
Y18059		Attachment site Corynephage 304L	Moreau, S. et al. "Analysis of the integration functions of φ304L: An integrase module among corynephages," Virology, 255(1):150-159 (1999)
Z21501	argS; lysA	Arginyl-tRNA synthetase; diaminopimelate decarboxylase (partial)	Oguiza, J.A. et al. "A gene encoding arginyl-tRNA synthetase is located in the upstream region of the lysA gene in Brevibacterium lactofermentum:
			Regulation of argS-lysA cluster expression by arginine," J. Bacteriol, 175(22):7356-7362 (1993)
Z21502	dapA; dapB	Dihydrodipicolinate synthase; dihydrodipicolinate reductase	Pisabarro, A. et al. "A cluster of three genes (dapA, orf2, and dapB) of Brevibacterium lactofermentum encodes dihydrodipicolinate reductase, and a
			third polypeptide of unknown function," J. Bacteriol., 175(9):2743-2749 (1993)
Z29563	thrC	Threonine synthase	Malumbres, M. et al. "Analysis and expression of the thrC gene of the encoded threonine synthase," Appl. Environ. Microbiol., 60(7)2209-2219 (1994)
Z46753	16S rDNA	Gene for 16S ribosomal RNA	
Z49822	sigA	SigA sigma factor	Oguiza, J.A. et al "Multiple sigma factor genes in Brevibacterium lactofermentum: Characterization of sigA and sigB," J. Bacteriol., 178(2):550-553 (1996)
Z49823	galE; dtxR	Catalytic activity UDP-galactose 4-	Oguiza, J.A. et al 'The galE gene encoding the UDP-galactose 4-epimerase of
		epimerase; diphtheria toxin regulatory protein	Brevibacterium lactofermentum is coupled transcriptionally to the dmdR gene," Gene, 177:103-107 (1996)
Z49824	orf1; sigB	?; SigB sigma factor	Oguiza, J.A. et al "Multiple sigma factor genes in Brevibacterium lactofermentum: Characterization of sigA and sigB," J. Bacteriol., 178(2):550-553 (1995)
Z66534		Transposase	Correia, A. et al. "Cloning and characterization of an IS-like element present in
			the genome of Brevibacterium lactofermentum ATCC 13869," Gene, 170(1):91-94 (1996)
	A A A A A A A A A A A A A A A A A A A		1 /0(1):91-94 (1990)

A sequence for this gene was published in the indicated reference. However, the sequence obtained by the inventors of the present application is significantly longer than the published version. It is believed that the published version relied on an incorrect start codon, and thus represents only a fragment of the actual coding region.

TABLE 3: Corynebacterium and Brevibacterium Strains Which May be Used in the Practice of the Invention

ලිනාගs	डिग्रेडब्रीस्ट 👉 🕒	WINGC	PRINT	Ferm  nrre  Ceci Neind  Gis  Ncrc Dswz	(दाष्ट्रा	NCIMIB	(C13)	NCTC	DSWZ
Brevibacterium	ammoniagenes	21054							
Brevibacterium	ammoniagenes	19350							
Brevibacterium	ammoniagenes	19351							
Brevibacterium	ammoniagenes	19352							
Brevibacterium	ammoniagenes	19353							
Brevibacterium	ammoniagenes	19354							
Brevibacterium	ammoniagenes	19355							
Brevibacterium	ammoniagenes	19356							
Brevibacterium	ammoniagenes	21055							
Brevibacterium	ammoniagenes	21077							
Brevibacterium	ammoniagenes	21553							
Brevibacterium	ammoniagenes	21580							
Brevibacterium	ammoniagenes	39101							
Brevibacterium	butanicum	21196							
Brevibacterium	divaricatum	21792	P928						
Brevibacterium	flavum	21474							
Brevibacterium	flavum	21129							:
Brevibacterium	flavum	21518							
Brevibacterium	flavum			B11474					
Brevibacterium	flavum			B11472					
Brevibacterium	flavum	21127							
Brevibacterium	flavum	21128							
Brevibacterium	flavum	21427							
Brevibacterium	flavum	21475							
Brevibacterium	flavum	21517							
Brevibacterium	flavum	21528							
Brevibacterium	flavum	21529							

Brevibacterium	flavum		B11477				
Brevibacterium	flavum		B11478	3			
Brevibacterium	flavum	21127					
Brevibacterium	flavum		B11474				
Brevibacterium	healii	15527					
Brevibacterium	ketoglutamicum	21004					
Brevibacterium	ketoglutamicum	21089					
Brevibacterium	ketosoreductum	21914					
Brevibacterium	lactofermentum			70			
Brevibacterium	lactofermentum			74			
Brevibacterium	lactofermentum			77			
Brevibacterium	lactofermentum	21798					
Brevibacterium	lactofermentum	21799					
Brevibacterium	lactofermentum	21800					
Brevibacterium	lactofermentum	21801					
Brevibacterium	lactofermentum		B11470	) [			
Brevibacterium	lactofermentum		B1147	1			
Brevibacterium	lactofermentum	21086					
Brevibacterium	lactofermentum	21420					
Brevibacterium	lactofermentum	21086					
Brevibacterium	lactofermentum	31269					
Brevibacterium	linens	9174					
Brevibacterium	linens	16261					
Brevibacterium	linens	8377					
Brevibacterium	paraffinolyticum				11160		
Brevibacterium	spec.					717.73	
Brevibacterium	spec.					717.73	
Brevibacterium	spec.	14604					
Brevibacterium	spec.	21860					·
Brevibacterium	spec.	21864					
Brevibacterium	spec.	21865	_				

Brevibacterium	spec.	21866		
Brevibacterium	spec.	19240		
Corynebacterium	acetoacidophilum	21476		
Corynebacterium	acetoacidophilum	13870		
Corynebacterium	acetoglutamicum		B11473	
Corynebacterium	acetoglutamicum		B11475	
Corynebacterium	acetoglutamicum	15806		
Corynebacterium	acetoglutamicum	21491		
Corynebacterium	acetoglutamicum	31270		
Corynebacterium	acetophilum		B3671	
Corynebacterium	ammoniagenes	, 6872		2399
Corynebacterium	ammoniagenes	15511		
Corynebacterium	fujiokense	21496		
Corynebacterium	glutamicum	14067		
Corynebacterium	glutamicum	39137		
Corynebacterium	glutamicum	21254		
Corynebacterium	glutamicum	21255		
Corynebacterium	glutamicum	31830		
Corynebacterium	glutamicum	13032		
Corynebacterium	glutamicum	14305		
Corynebacterium	glutamicum	15455		
Corynebacterium	glutamicum	13058		
Corynebacterium	glutamicum	13059		•
Corynebacterium	glutamicum	13060		
Corynebacterium	glutamicum	21492		
Corynebacterium	glutamicum	21513		
Corynebacterium	glutamicum	21526		
Corynebacterium	glutamicum	21543		
Corynebacterium	glutamicum	13287		
Corynebacterium	glutamicum	21851		
Corvnebacterium	glutamicum	21253		_

Corynebacterium	glutamicum	21514		
Corynebacterium	glutamicum	21516		
Corynebacterium	glutamicum	21299		
Corynebacterium	glutamicum	21300		
Corynebacterium	glutamicum	39684		
Corynebacterium	glutamicum	21488		
Corynebacterium	glutamicum	21649		
Corynebacterium	glutamicum	21650		
Corynebacterium	glutamicum	19223		
Corynebacterium	glutamicum	13869		
Corynebacterium	glutamicum	21157		_
Corynebacterium	glutamicum	21158		
Corynebacterium	glutamicum	21159		
Corynebacterium	glutamicum	21355		
Corynebacterium	glutamicum	31808		
Corynebacterium	glutamicum	21674		
Corynebacterium	glutamicum	21562		
Corynebacterium	glutamicum	21563		
Corynebacterium	glutamicum	21564		
Corynebacterium	glutamicum	21565		
Corynebacterium	glutamicum	21566		
Corynebacterium	glutamicum	21567		
Corynebacterium	glutamicum	21568		
Corynebacterium	glutamicum	21569	,	
Corynebacterium	glutamicum	21570		
Corynebacterium	glutamicum	21571		
Corynebacterium	glutamicum	21572		
Corynebacterium	glutamicum	21573		
Corynebacterium	glutamicum	21579		
Corynebacterium	glutamicum	19049		
Corynebacterium	glutamicum	19050		

Corynebacterium	glutamicum	19051					
Corynebacterium	glutamicum	19052					
Corynebacterium	glutamicum	19053					
Corynebacterium	glutamicum	19054					
Corynebacterium	glutamicum	19055					
Corynebacterium	glutamicum	19056				-	
Corynebacterium	glutamicum	19057					
Corynebacterium	glutamicum	19058					
Corynebacterium	glutamicum	19059					
Corynebacterium	glutamicum	19060					
Corynebacterium	glutamicum	19185					
Corynebacterium	glutamicum	13286					
Corynebacterium	glutamicum	21515					
Corynebacterium	glutamicum	21527					
Corynebacterium	glutamicum	21544					
Corynebacterium	glutamicum	21492		,			
Corynebacterium	glutamicum			B8183			
Corynebacterium	glutamicum			B8182			
Corynebacterium	glutamicum			B12416			
Corynebacterium	glutamicum			B12417			
Corynebacterium	glutamicum			B12418			
Corynebacterium	glutamicum			B11476			
Corynebacterium	glutamicum	21608					
Corynebacterium	lilium		P973				
Corynebacterium	nitrilophilus	21419			11594		
Corynebacterium	spec.		P4445				
Corynebacterium	spec.		P4446				
Corynebacterium	spec.	31088					
Corynebacterium	spec.	31089					
Corynebacterium	spec.	31090					
Corynebacterium	spec.	31090					

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Corynebacterium spec.	spec.	31090	
Corynebacterium spec.	spec.	15954	20145
Corynebacterium	spec.	21857	
Corynebacterium spec.	spec.	21862	
Corynebacterium	ı spec.	21863	

ATCC: American Type Culture Collection, Rockville, MD, USA

FERM: Fermentation Research Institute, Chiba, Japan

NRRL: ARS Culture Collection, Northern Regional Research Laboratory, Peoria, IL, USA

CECT: Coleccion Espanola de Cultivos Tipo, Valencia, Spain

NCIMB: National Collection of Industrial and Marine Bacteria Ltd., Aberdeen, UK

CBS: Centraalbureau voor Schimmelcultures, Baarn, NL

NCTC: National Collection of Type Cultures, London, UK

DSMZ: Deutsche Sammlung von Mikroorganismen und Zellkulturen, Braunschweig, Germany

For reference see Sugawara, H. et al. (1993) World directory of collections of cultures of microorganisms: Bacteria, fungi and yeasts (4th edn), World federation for culture collections world data center on microorganisms, Saimata, Japen.

## TABLE 4: ALIGNMENT RESULTS

# Q	length (NT)	length Genbank Hit (NT)	Length		Accession Name of Genbank Hit	Source of Genbank Hit	% homology (GAP)	Date of Deposit
rxa00009 1023	1023	GB_IN1:CELZK563 GB_IN1:CELZK563	29655 29655	U40061 U40061	Caenorhabditis elegans cosmid ZK563. Caenorhabditis elegans cosmid ZK563.	Caenorhabditis elegans Caenorhabditis elegans	33,694 36,040	9-Nov-95 9-Nov-95
та00010	810	GB_BA1:MTCY164 3 GB_BA1:MTFTSX 4 GB_BA1:SHGCPIR 1	39150 4068 107379	Z95150 X70031 X86780	Mycobacterium tuberculosis H37Rv complete genome; segment 135/162. Mycobacterium tuberculosis M.tuberculosis fisX and fisE (partial) genes. S.hygroscopicus gene cluster for polyketide immunosuppressant rapamycin. Streptomyces hygroscopicus		38,442 63,158 38,875	19-Jun-98 06-MAR-1997 16-Aug-96
гха00024 1068	1068		й 10000 й 10000		Caenorhabditis elegans chromosome V clone Y113G7, *** SEQUENCING IN PROGRESS ***, in unordered pieces. Caenorhabditis elegans chromosome V clone Y113G7, *** SEQUENCING IN PROGRESS ***, in unordered pieces.	Caenorhabditis elegans Caenorhabditis elegans	36,217 36,217 35,217	12-Jan-99 12-Jan-99
rxa00048 1782	1782	GB_HTG3:AC008905 1	129915 129915	AC008905 AC008905	Arabidopsis trialiaria DNA cirioritosome 4, DAC done FIGIZ (ESSA project).  Homo sapiens chromosome 5 clone CITB-H1_2259114, *** SEQUENCING IN PROGRESS ***, 40 unordered pieces.  Homo sapiens chromosome 5 clone CITB-H1_2259114, *** SEQUENCING IN PROGRESC ***	Homo sapiens Homo sapiens	38,826 38,826	3-Aug-99
rxa00070	555	GB_HTG3:AC008905 CB_BA2:BPEFUR	129915	AC008905 L31851	IN PRUCKESS ***, 40 unordered pieces.  Homo sapiens chromosome 5 clone CITB-H1_2259114, *** SEQUENCING IN PROGRESS ***, 40 unordered pieces.  Bordetella pertussis DNA repair protein (recN) gene, partial cds; iron requilatory protein (fur) gene, complete cds.	Homo sapiens Bordetella pertussis	37,379 45,756	3-Aug-99 17-Apr-95
rxa00078	537	GB_BA2:BPU11699 537 GB_BA1:BTFURRECN 1106 GB_PR3:HUMCOL2A1Z31001 GB_HTG2:AC006721 13555	537 1106 231001 135550	U11699 Z48227 L10347 AC006721	Bordetella pertussis ferric uptake regulator (fur) gene, complete cds.  B.pertussis fur gene for ferric uptake regulator (fur) gene, complete cds.  Human pro-alpha1 type II collagen (COL2A1) gene exons 1-54, complete cds.  Caenorhabditis elegans clone Y18H1, *** SEQUENCING IN PROGRESS	Bordetella pertussis Bordetella pertussis Homo sapiens Caenorhabditis elegans	47,119 45,756 39,010 40,661	14-Jan-95 10-Feb-95 3-Aug-95 23-Feb-99
rxa00088	668	GB_RO:MMCGT6	135550 3009 3009	AC006721 U48896 U48896	T., 5 unordered pleces. Caenorhabditis elegans clone Y18H1, *** SEQUENCING IN PROGRESS ***, 5 unordered pieces. Mus musculus UDP-galactose:ceramide galactosyltransferase (Cgt) gene, exon 6 and complete cds. Mus musculus UDP-galactose:ceramide galactosyltransferase (Cgt) gene, exon 6 and complete cds.	Caenorhabditis elegans Mus musculus Mus musculus	40,661 35,455 34,439	23-Feb-99 1-Nov-96 1-Nov-96
ка00100 723 ка00135 1377 жа00143 1605	723 1377 1605	GB_PL1:CAC41C10 3887 GB_PR4:AC007115 1808 GB_PR4:AC007115 1808 GB_BA1:MTCY373 3551 GB_BA1:MLU15186 3624 GB_BA1:MTMURAGEN1257 GB_PAT:I92051 1107	38874 180821 180821 35516 36241 1257 1107	AL033501 AC007115 AC007115 Z73419 U15186 X96711	C.albicans cosmid Ca41C10.  Homo sapiens chromosome 12 clone 917O5, complete sequence.  Homo sapiens chromosome 12 clone 917O5, complete sequence.  Mycobacterium tuberculosis H37Rv complete genome; segment 57/162.  Mycobacterium leprae cosmid L471.  M.tuberculosis murA gene.  Sequence 18 from patent US 5726299.	Candida albicans Homo sapiens Homo sapiens Mycobacterium tuberculosis Mycobacterium teprae Mycobacterium tuberculosis Unknown.	36,222 33,050 34,993 60,639 38,377 61,575 37,773	10-Nov-98 17-Aug-99 17-Aug-99 17-Jun-98 09-MAR-1995 22-MAR-1996 01-DEC-1998

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	GB_PAT:178761		178761	Sequence 17 from patent US 5693781.		37,773	3-Apr-98
~200177 1101	GB_BA1:MTCY28	40163	Z95890	Mycobacterium tuberculosis H37Rv complete genome; segment 79/162.  PDCI-11-365I 6 TV PDCI-11 Home senions genomic clone RDCI-11-365I 6	Mycobacterium tuberculosis Homo saniens	36,984 38,551	18-Jun-98 19-MAY-1999
	GB_PL2:AF017646	3394	AF017646	Schizosaccharomyces pombe TFIIH subunit p47 (tfh47) gene, complete	Schizosaccharomyces	38,122	17-MAR-1999
	GB PI 1:SPCC1682	37404	AI 031525	cds. S nombe chromosome III cosmid c1682.	pombe Schizosaccharomyces	33.983	14-DEC-1998
					pombe	<u>.</u>	
rxa00178 1008	38 GB_BA1:AB016258	2260	AB016258	Arthrobacter sp. gene for maleylacetate reductase and hydroxyquinol 1,2-	Arthrobacter sp.	65,182	8-Sep-99
	GB_BA1:CGPUTP	3791	Y09163	droxygeriase, parta and comprete cos. C.glutamicum putP gene.	Corynebacterium glutamicum 38,806	38,806	8-Sep-97
			G05495	human STS WI-5918.	Homo sapiens	39,925	ca-unc-s
rxa00277 1684	34 GB_BA1:MTCY22G10	10 35420	Z84724	Mycobacterium tuberculosis H37Rv complete genome; segment 21/162.	Mycobacterium tuberculosis	39,976	17-Jun-98
	GB_IN1:CELT03F1	38643	<b>U88169</b>	Caenorhabditis elegans cosmid T03F1.	Caenorhabditis elegans	35,127	7-Feb-97
	GB_IN2:CELK02A2	38261	U23171	Caenorhabditis elegans cosmid K02A2.	Caenorhabditis elegans	36,166	21-MAY-1999
rxa00372 1200	00 GB_IN2:AC005452	79333	AC005452	Drosophila melanogaster, chromosome 2R, region 43B2-43C2, P1 clone	Drosophila melanogaster	37,006	26-Nov-98
				DS07185, complete sequence.			
	GB_IN2:AC005452	79333	AC005452	Drosophila melanogaster, chromosome 2R, region 43B2-43C2, P1 clone	Drosophila melanogaster	34,907	26-Nov-98
				DS07185, complete sequence.	:		
	GB_IN1:CELW03F8		AF039041	Caenorhabditis elegans cosmid W03F8.	Caenorhabditis elegans	40,712	1-Jan-98
rxa00389 1683		772	AB010703	Thelleria sp. gene for major piroplasm surface protein, partial cds, isolate	I nellerla sp.	40,285	98-JdY-91
	GB BA1:11108911	619	1108911	Kampnaeng Saen. Lactobacillus leichmannii outative D-alanine:D-alanine lioase (ddl) gene.	Lactobacillus leichmannii	40.194	16-Feb-96
		•		partial cds.			
	GB IN1:TPMS1	822	Z48740	T.parva Tpms1 gene for merozoite surface glycoprotein.	Theileria parva	38,902	15-MAY-1995
rxa00467 792				Homo sapiens jun dimerization protein gene, partial cds; cfos gene,	Homo sapiens	37,995	7-Apr-99
				complete cds; and unknown gene.			
	GB_PR4:DJ293M10	202267	AF111167	Homo sapiens jun dimerization protein gene, partial cds; cfos gene,	Homo sapiens	36,639	7-Apr-99
				complete cds; and unknown gene.			
	GB_IN1:CEW01C9		Z49969	Caenorhabditis elegans cosmid W01C9, complete sequence.	Caenorhabditis elegans	37,980	23-Nov-98
rxa00499 1404		42732	AC007206	Homo sapiens chromosome 19, cosmid R27370, complete sequence.	Homo sapiens	34,982	4-Apr-99
	GB_EST26:Al344735	5 462	AI344735	qp05a10.x1 NCI_CGAP_Kid5 Homo sapiens cDNA clone IMAGE:1917114 3' similar to ob:M15800 T-I YMPHOCYTE MATURATION-ASSOCIATED	Homo sapiens	42,675	2-Feb-99
				PROTEIN (HUMAN);, mRNA sequence.			
	GB_PR4:AC006479	161837	AC006479	Homo sapiens clone DJ1051J04, complete sequence.	Homo sapiens	38,462	11-Nov-99
rxa00508 1206		1 84245	AC007111	Homo sapiens chromosome 16 clone 1-8F, *** SEQUENCING IN	Homo sapiens	37,931	18-MAR-1999
	CB UTC2:AC007111	1 842AE	AC007111	HOUSE Springs chromosome 16 clone 1-8E *** SECTIFICATIONS IN	Homo sapiens	37 931	18-MAR-1999
	11 7000 Y. 7000 T. 7000			PROGRESS *** 2 ordered pieces.		5	
	GB_VI:AF141890	1791	AF141890	Columbid herpesvirus 1 DNA-dependent DNA polymerase gene, partial cds. columbid herpesvirus 1	columbid herpesvirus 1	39,401	7-Jul-99
rxa00569 1149	49 GB_PAT:115213 GB_PAT:E07353	3728 3728	115213 E07353	Sequence 1 from patent US 5460951. cDNA encoding bone-related carboxypeptidase-like protein, OSF-5.	Unknown. Mus sp.	41,244 41,244	2-Apr-96 29-Sep-97

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		GB_HTG1:CEY70G10 152184 AL020987	152184	AL020987	Caenorhabditis elegans chromosome III clone Y70G10, *** SEQUENCING C	Caenorhabditis elegans	34,148	12-DEC-1997
rxa006,12 1077		GB_HTG2:AC005020	177756	AC005020	Home sapiens clone GS259H13, *** SEQUENCING IN PROGRESS ***, 4	Homo sapiens	34,551	12-Jun-98
	_	GB_HTG2:AC005020	177756	AC005020	unordered pieces.  Homo sapiens clone GS259H13, *** SEQUENCING IN PROGRESS ***, 4	Homo sapiens	34,551	12-Jun-98
	-	GB_HTG2:AC005020	177756	AC005020	undurered pieces.  Homo sapiens clone GS259H13, *** SEQUENCING IN PROGRESS ***, 4	Homo sapiens	37,628	12-Jun-98
rxa00615 705		GB_GSS15:AQ622921	517	AQ622921	unordered pieces.  HS_5351_A1_A08_T7A RPCI-11 Human Male BAC Library Homo sapiens  annumic clone Plate=927 Col=15 Row=A genomic survey sequence.	Homo sapiens	38,254	16-Jun-99
		GB_GSS3:B36703	432	B36703	0 >-	Homo sapiens	44,981	17-OCT-1997
		GB_EST25:Al245926	572	Al245926	sequence. qk33c08.x1 NCI_CGAP_Co8 Homo sapiens cDNA clone IMAGE:1870766 F 3' similar to SW:COPG_BOVIN P53620 COATOMER GAMMA SUBUNIT;,	Homo sapiens	38,902	28-Jan-99
rxa00621 906		GB_EST1:D36491	360	D36491	ji Kohara unpublished cDNA Caenorhabditis elegans	Caenorhabditis elegans	40,390	8-Aug-94
		GB_IN2:CELC16A3 GB_HTG3:AC009311	34968 160198	U41534 AC009311	Caenorhabditis elegans cosmid C16A3. Homo sapiens clone NH0311L03, *** SEQUENCING IN PROGRESS ***, 3	Caenorhabditis elegans Homo sapiens	35,477 38,636	18-MAY-1999 13-Aug-99
rxa00622 15	1539	GB_BA1:AB004795 GB_BA1:MBOPII GB_IN2:AF078916	3039 2392 2960	AB004795 D38405 AF078916	Pseudomonas sp. gene for dipeptidyl aminopeptidase, complete cds. Moraxella lacunata gene for protease II, complete cds. Trypanosoma brucei brucei oligopeptidase B (opb) gene, complete cds.	Pseudomonas sp. Moraxella lacunata Trypanosoma brucei	54,721 50,167 48,076	5-Feb-99 8-Feb-99 08-OCT-1999
rxa00639 978		GB_BA2:AF043741 GB_BA1:D83237	1223	AF043741 D83237	Rhodococcus rhodochrous catechol 1,2-dioxygenase (catA) gene, complete cds.  Rhodococcus ervthropolis DNA for catechol 1,2-dioxgenase, complete cds.		66,940 65,440	27-Aug-98 1-Sep-99
7200641 ואפס		GB_BA2:AF134348	7224 5000	X99622. AF134348	Rhodococcus opacus catR, catA, catB, catC genes and five ORFs. Pseudomonas putida plasmid pDK1 toluate 1,2 dioxygenase subunit (xylX), toluate 1,2 dioxygenase subunit (xylY), and toluate 1,2 dioxygenase subunit (xylZ) genes, complete cds, and 1,2-dihydroxycyclohexa-3,5-diene carboxylate dehydrogenase (xylL) gene, partial cds.		63,617 59,863	24-Sep-97 20-MAY-1999
ка00642 615		GB_BA1:PWWXYL GB_BA1:PCCBDABC GB_BA2:AF134348	9037 3548 5000	M64747 X79076 AF134348	Pseudomonas putida plasmid pWW0 meta operon, 5' genes. P.cepacia (2CBS) cbdA, cbdB and cbdC genes. Pseudomonas putida plasmid pDK1 toluate 1,2 dioxygenase subunit (xylX), toluate 1,2 dioxygenase subunit (xylY), and toluate 1,2 dioxygenase subunit (xylZ) genes, complete cds; and 1,2-dihydroxycyclohexa-3,5-diene carboxylate dehydrogenase (xylL) gene, partial cds.	Plasmid pVWV0 Burkholderia cepacia Pseudomonas putida	59,588 55,410 60,920	26-Apr-93 3-Apr-97 20-MAY-1999
		GB_BA1:PWWXYL GB_GSS11:AQ274007	9037 637	M64747 AQ274007	Pseudomonas putida plasmid pWW0 meta operon, 5' genes. nbxb0032I07f CUGI Rice BAC Library Oryza sativa genomic clone nbxb0032I07f, genomic survey sequence.	Plasmid pWW0 Oryza sativa	58,756 41,390	26-Apr-93 3-Nov-98

a 53,871 20-MAY-1999	52,603 26-Apr-93 43,865 16-Jun-98	55,626 26-Apr-93 a 50,410 20-MAY-1999	40,138 28-Aug-98	40,636 6-Aug-97	38,406 6-Aug-97	arcutosis 57,976 24-Jun-99 ae 39,669 27-Aug-99 36,154 05-MAY-1999	color 36,836 15-Jan-99 42,027 01-OCT-1998	laster 35,531 27-OCT-1999	37,295 03-DEC-1999	37,295 03-DEC-1999	35,650 21-Aug-97	2 18- hil-98
, Pseudomonas putida t	Plasmid pWW0 Mus musculus	Plasmid pWW0 , Pseudomonas putida t	Homo sapiens	Oryza sativa	Oryza sativa	Mycobacterium tuberculosis Mycobacterium leprae , Homo sapiens	Streptomyces coelicolor Homo sapiens	Drosophila melanogaster	Homo sapiens	Homo sapiens	Homo sapiens	Pseudomonas aeru
Pseudomonas putida plasmid pDK1 toluate 1,2 dioxygenase subunit (xylK), toluate 1,2 dioxygenase subunit (xylY), and toluate 1,2 dioxygenase subunit (xylZ) genes, complete cds; and 1,2-dihydroxycyclohexa-3,5-diene carboxylate dehydrogenase (xylL) gene, partial cds.	Pseudomonas putida plasmid pWWV0 meta operon, 5' genes. ua97f07.r1 Soares mouse mammary gland NbMMG Mus musculus cDNA clone IMAGE:1365445 5' similar to SW:DUS7_RAT Q63340 DUAL			c26090 Rice callus cDNA Oryza sativa cDNA clone C11617_1A, mRNA	sequence. C26090 Rice callus cDNA Oryza sativa cDNA clone C11617_1A, mRNA sequence.		<ul> <li>b unordered pieces.</li> <li>Streptomyces coelicolor cosmid 3C8.</li> <li>i Homo sapiens chromosome 16, BAC clone 462G18 (LANL), complete</li> </ul>	sequence.  Drosophila melanogaster, chromosome 2L, region 38A5-38B4, BAC clone BACR48M05, complete sequence.			SECUENCING IN PROCEEDS In unitation process.  Human chromosome 12p15 BAC clone CIT987SK-99D8 complete	Sequence.   Desurfaments seguriness segurations & francturase Desertà (nesertà) nans. Desurfaments segurinines
AF134348	M64747 A1020666	M64747 AF134348	A1038396	C26090	C26090	AL021897 AL049491 AC007482	AL023861 AC005736	AC005719	AL109942	AL109942	U91327	AF010184
2000	9037 328	9037 5000	438	414	414	67200 34714 155357	33095 215441	188357	203460	203460	129252	1494
GB_BA2:AF134348	GB_BA1:PWWXYL GB_EST22:Al020666	GB_BA1:PWWXYL GB_BA2:AF134348	GB_EST22:A1038396	GB_EST16:C26090	GB_EST16:C26090	GB_BA1:MTV017 GB_BA1:MLCB1222 GB_HTG2:AC007482	GB_BA1:SC3C8 GB_PR3:AC005736	GB_IN2:AC005719	GB_HTG2:HSJ473J16 203460 AL109942	GB_HTG2:HSJ473J16	GB_PR2:HSU91327	GR BA2.AF010184
ка00643 1659		ка00644 951		rxa00658 816		rxa00663 1497	rxa00675 915		rxa00762 999			929 CZ2002X

Table 4, Page 4

01-DEC-1998

3-Apr-98

92,701 92,701 47,791 35,536 38,006

Mycobacterium tuberculosis

Unknown. Unknown.

Synechocystis sp. Synechocystis sp.

M.tuberculosis PstS-2 gene. Synechocystis sp. PCC6803 complete genome, 9/27, 1056467-1188885. Synechocystis sp. PCC6803 complete genome, 9/27, 1056467-1188885.

Sequence 10 from patent US 5726299. Sequence 10 from patent US 5693781.

Z48056 D90907

GB\_PAT:192043
GB\_PAT:178754
GB\_BAT:MTPST2GN
GB\_BA1:D90907
GB\_BA1:D90907

rxa00778 1248

1347 132419 132419

192043 178754

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complete cds.

24-Apr-99 7-Feb-99 7-Feb-99

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rxa00787 2025	2025	GB_PL1:SCX11RA	36849	X91258	risiae DNA from chromosome XII right arm including ACE2, CKI1,	Saccharomyces cerevisiae	36,122	13-OCT-1995
							7	1007
		GB_PL2:YSCL9606 GB_PL1:SCX11RA	29154 36849	U53881 X91258	Saccharomyces cerevisiae chromosome XII cosmid 9606. S.cerevisiae DNA from chromosome XII right arm including ACE2, CKI1,	Saccharomyces cerevisiae Saccharomyces cerevisiae	36,122 37,198	25-OC1-1997 13-OCT-1995
		ì						
rxa00792 1320	1320	GB_PR4:AC004841	132072	AC004841	Homo sapiens PAC clone DJ0607J23 from 7q21.2-q31.1, complete	Homo sapiens	37,452	18-MAR-1999
		GB_HTG2:AC006706	180664	AC006706	Sequence.  Caenorhabditis elegans clone Y110A2, *** SEQUENCING IN PROGRESS  *** 4 unordered pieces.	Caenorhabditis elegans	34,824	23-Feb-99
		GB_HTG2:AC006706	180664	AC006706	Caenorhabditis elegans clone Y110A2, *** SEQUENCING IN PROGRESS ***, 4 unordered pieces.	Caenorhabditis elegans	34,824	23-Feb-99
rxa00857 1313	1313	GB_BA1:MTV002	56414	AL008967	osis H37Rv complete genome; segment 122/162.	Mycobacterium tuberculosis	38,080	17-Jun-98
		GB_BA1:MSGY154	40221	AD000002	Mycobacterium tuberculosis sequence from clone y154.	Mycobacterium tuberculosis	68,345	03-DEC-1996
		GB_BA1:MLCB33	42224	Z94723		Mycobacterium leprae	38,824	24-Jun-97
rxa00877 17	1788	GB_PAT:192050	267	192050	Sequence 17 from patent US 5726299.	Unknown.	62,787	01-DEC-1998
		GB_PAT:178760	292	178760		Unknown.	62,787	3-Apr-98
		GB_BA2:AE000426	10240	AE000426	Escherichia coli K-12 MG1655 section 316 of 400 of the complete genome.	Escherichia coli	36,456	12-Nov-98
rxa00888 1140		GB_BA1:MTCY27	27548	Z95208	Mycobacterium tuberculosis H37Rv complete genome; segment 104/162.	Mycobacterium tuberculosis	40,165	17-Jun-98
		GB_BA1:U00016	42931	U00016	Mycobacterium leprae cosmid B1937.	Mycobacterium leprae	58,444	01-MAR-1994
		GB_BA1:ECU82598	136742	U82598	f minutes 9 to 12.	Escherichia coli	37,876	15-Jan-97
rxa00892 10	1017	GB_BA2:AE000817	13157	AE000817	Methanobacterium thermoautotrophicum from bases 251486 to 264642	Methanobacterium	36,710	15-Nov-97
					-	thermoautotrophicum		
		GB_EST29:AI620549	239	AI620549	3773	Homo sapiens	38,075	21-Apr-99
					<ol> <li>similar to gb:X60708_rna1 DIPEPTIDYL PEPTIDASE IV (HUMAN);, mRNA sequence.</li> </ol>			
		GB_BA2:AE000817	13157	AE000817	Methanobacterium thermoautotrophicum from bases 251486 to 264642	Methanobacterium	35,650	15-Nov-97
		•				thermoautotrophicum		
rxa00897 1	1128	GB_PR3:HS246D7	28011	AL031843	ie 22q13.1-13.33.	Homo sapiens	38,724	23-Nov-99
		GB PR3:HSDJ185D5	24387	AL118498	Contains ESTS, a GSS and an STS, complete sequence. Human DNA sequence from clone 185D5 on chromosome 22, complete	Homo sapiens	37,021	23-Nov-99
		ı						
		GB_PR3:HS246D7	28011	AL031843	Human DNA sequence from clone 246D7 on chromosome 22q13.1-13.33.	Homo sapiens	36,054	23-Nov-99
rxa00944 1095	1095	GB_BA1:ECU68759	1531	U68759	e (onr) gene,	Enterobacter cloacae	43,041	14-DEC-1996
		ı			complete cds.			
		GB_PAT:A59288	1531	A59288	Sequence 1 from Patent WO9703201.	unidentified	43,041	06-MAR-1998
		GB_EST23:A1099394	601	A1099394		Mus musculus	37,225	20-Aug-98
					IMAGE:1482040 5' similar to gb:U21301 Mus musculus c-mer tyrosine			
,	9				ige.		0	
rxa00964 1248	1248	GB_HTG6:AC009794	152794	AC009794	Homo sapiens chromosome 4 clone RP11-343C10 map 4, *** SEQUENCING IN PROGRESS *** 33 unordered pieces.	Homo sapiens	34,762	03-DEC-1999
		GB_HTG6:AC009794	152794	152794 AC009794	sapiens chromosome 4 clone RP11-343C10 map 4, ***	Homo sapiens	35,708	03-DEC-1999
					SECUENCING IN PROGRESS ***, 33 unordered pieces.			

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				TABLE 4. ALIGNATURE ALEGALIS			
rxa00982 1629	629	GB_BA1:BLARGS 2501	I Z21501	B.lactofermentum argS and lysA genes for arginyl-tRNA synthetase and diaminonimalste departments.	Corynebacterium glutamicum 39,003	39,003	28-DEC-1993
		GB_BA1:CGXLYSA 2344	t X54740	Corynebacterium glutamicum argS-lysA operon gene for the upstream region of the arginyl-tRNA synthetase and diaminopimelate decarboxylase (EC 4.1.1.20).	Corynebacterium glutamicum 41,435	41,435	30-Jun-93
		GB_PAT:E14508 3579	9 E14508	ONA encoding Brevibacterium diaminopimelic acid decarboxylase and arginyl-tRNA synthase.	Corynebacterium glutamicum 40,566	40,566	28-Jul-99
rxa01014 2.	2724	GB_BA1:STMAMPEPN 2849 GB_BA1:SC7H2 42655	33 AL021246 3 L23172 55 AL109732	ilosis H37Rv complete genome; segment 108/162. aminopeptidase N gene, complete cds. ir cosmid 7H2.	Mycobacterium tuberculosis 5 Streptomyces lividans 5 Streptomyces coelicolor 3 A3(2)	56,167 57,067 37,551	17-Jun-98 18-MAY-1994 2-Aug-99
rxa01022 1;	1203	GB_PAT:A68384 1080 GB_BA2:AF077728 1346	A68384 AF077728	Sequence 1 from Patent WO9748809. Mycobacterium smegmatis D-alanine:D-alanine ligase gene, complete cds.	bacterium avium bacterium smegmatis	56,913 57,203	06-MAY-1999 1-Jan-99
rxa01055		GB_BA1:MSGB1723CS38477	77 L78825	Mycobacterium leprae cosmid B1723 DNA sequence.	Mycobacterium leprae 5	54,599	15-Jun-96
rxa01056 10	1023	GB_BA2:AE001715 11086 GB_EST38:AW046857 161	36 AE001715 AW046857	Thermotoga maritima section 27 of 136 of the complete genome. UI-M-BH1-akI-a-04-0-UI.s1 NIH_BMAP_M_S2 Mus musculus cDNA clone	Thermotoga maritima 3 Mus musculus 4	39,034 45,963	2-Jun-99 18-Sep-99
		GB_EST38:AW049435 244	AW049435			40,984	18-Sep-99
rxa01057 16	1626	GB_PL1:LPAJ5046 656 GB_PL2:SPAC806 22870	AJ225046 70 AL117212		eruvianum omyces	37,117 38,211	22-Jul-98 24-Nov-99
		GB_PL2:SPAC806 22870	70 AL117212	S.pombe chromosome I cosmid c806.	accharomyces	36,934	24-Nov-99
rxa01082 7	783	GB_BA2:AF112535 4363	3 AF112535	Corynebacterium glutamicum putative glutaredoxin NrdH (nrdH), NrdI (nrdI), and ribonucleotide reductase albha-chain (nrdE) genes, complete cds.	pombe Corynebacterium glutamicum 99,794	99,794	5-Aug-99
rxa01113_2	260	GB_PL2:TAE237897 8020 GB_PL2:AF076680 10499 GB_VI:ASU02468 11424	AJ237897 99 AF076680 24 U02468		Triticum aestivum Aegilops tauschii African swine fever virus	37,132 38,651 31,923	1-Nov-99 14-MAY-1999 28-Apr-94
		GB_VI:ASU18466 170101 GB_GSS5:AQ752779 1647	101 U18466 7 AQ752779	A506R, and A542R) genes, complete cds.  African swine fever virus, complete genome.  HS_5569_B1_D02_SP6 RPCI-11 Human Male BAC Library Homo sapiens  Approprie close Dista=1446 Col=3 Row=H genomic survey contents.	African swine fever virus 3 Homo sapiens 3	31,923 37,154	22-Apr-95 19-Jul-99
rxa01115 8	876					40,850	26-DEC-1998
rxa01116 7;	735					37,326 35,877 40,616 64,099	10-Sep-96 14-Jan-98 21-Jun-99 06-DEC-1999
		GB_BA1:MTCY07A7 23967	37 Z95556	Mycobacterium tuberculosis H37Rv complete genome; segment 109/162.	Mycobacterium tuberculosis 4	41,716	17-Jun-98

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				IABLE 4: ALIGINIVENT RESOLUS			
rxa01117 864	GB_BA2:AF109386	6551	AF109386	Streptomyces sp. 2065 protocatechuaic acid catabolic gene cluster, complete sequence:	Streptomyces sp. 2065	62,116	06-DEC-1999
	GB_BA2:AF003947	5475	AF003947	is succinyl CoA:3-oxoadipate CoA transferase subunit e, partial cds, protocatechuate dioxygenase beta ocatechuate dioxygenase alpha subunit (pcaG), 3-onate cycloisomerase homolog (pcaB), 3-oxoadipate sel4-carboxymuconolactone decarboxylase (pcaL) and complete cds, and 3-oxoadipyl CoA thiolase homolog	Rhodococcus opacus	36,712	12-MAR-1998
	GB BA1:XCLPSIJ	2578	Y11313	X.campestris lpsl, lpsJ, xanA genes and orfX.	Xanthomonas campestris	39,833	20-Jan-98
1401	GB BA1-MTV008	63033	AI 021246	Mycobacterium tuberculosis H37Ry complete genome: segment 108/162.	Mycobacterium tuberculosis	36,715	17-Jun-98
1041 071 10841	GB_BA1:CA.110321	6710	A.1010321	Caulobacter crescentus partial tig gene and cloP, cicA, cloX, lon genes.		63,311	01-OCT-1998
	GB_BA2:AF150957	4440	AF150957	Azospirillum brasilense trigger factor (tig), heat-shock protein ClpP (clpP), and heat-shock protein ClpX (clpX) genes, complete cds; and Lon protease (lon) gene partial cds.		60,613	7-Jun-99
2201126 E83	GB HTC3-AC000100	86408	AC009199	(1911) gene, parital 325. Drosophila melanogaster chomosome 2 clone BACR10.123 (D1024) RPCI. Drosophila melanogaster		35,294	20-Sep-99
		3		98 10.J.23 map 37B-37B strain y; cn bw sp, *** SEQUENCING IN PROGRESS ***, 79 unordered pieces.			
	GB_HTG3:AC009199	66498	AC009199	Drosophila melanogaster chromosome 2 clone BACR10J23 (D1024) RPCI- Drosophila melanogaster		35,294	20-Sep-99
				98 10.J.23 map 378-37B strain y; cn bw sp, *** SEQUENCING IN PROGRESS ***, 79 unordered pieces.			
	GB_PL1:AB016880	81284	AB016880	Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone: MTG10, complete sequence.		34,477	20-Nov-99
rxa01181 980	GB BA1:MLCB22	40281	Z98741	Mycobacterium leprae cosmid B22.	Mycobacterium leprae	61,570	22-Aug-97
	GB_BA1:MTCY190	34150	Z70283	Mycobacterium tuberculosis H37Rv complete genome; segment 98/162.	Mycobacterium tuberculosis	60,434	17-Jun-98
	GB_BA1:SC5F7	40024	AL096872	Streptomyces coelicolor cosmid 5F7.	Streptomyces coelicolor A3(2)	57,011	22-Jul-99
rxa01236 1068	GB_EST3:H01832	381	H01832	yj28c11.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:150068 3', mRNA sequence.	Homo sapiens	41,406	19-Jun-95
	GR PR4-AC004850	105891	AC004850	Homo sapiens PAC clone DJ0665C04 from 7p14-p13, complete sequence.	Homo sapiens	37,428	26-Feb-99
	GB_GSS11:AQ304150		AQ304150	HS_3208_A1_D12_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3208 Col=23 Row=G, genomic survey		37,421	16-DEC-1998
2201284 1302	300/TMT.008	121125	AI 022121	sequence. Mycobacterium tuberculosis H37Ry complete genome: segment 155/162	Mycobacterium tuberculosis	58.315	24-Jun-99
17401204 1332		02777820	101263	M Jense genomic das seguence cosmid 5577			14-Jun-96
	071/0007/10000/1/00 071/0007/1/000000	25645	A1 023506	Michael genomic and sequence, cosmic con	Mycobacterium leprae	37 645	27-Aug-99
9704 0701000			Y01182	Mycobacterium reprae cosmila 02407. Recterial so padrial 168 rRNA pepe (clone group (310)	injection in the community of the commun	41,228	15-Jul-96
	GR BA1-BSP.IN12D		Z69277	Bacterial sp. partial 16S rRNA gene (clone group JN12d).	Bacteria	38,905	24-Jun-98
	GB_EST7:W93397		W93397	2d95b03.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone	Homo sapiens	40,516	25-Nov-96
rxa01277 2127	GB_PL2:AF111709	52684	AF111709	Oryza sativa subsp. indica Retrosat 1 retrotransposon and Ty3-Gypsy type Retrosat 2 retrotransposon, complete sequences; and unknown genes.	Oryza sativa subsp. indica	37,410	26-Apr-99
	GB_IN1:CELZC250	34372	AF003383	Caenorhabditis elegans cosmid ZC250.	Caenorhabditis elegans	35,506	14-MAY-1997

				TABLE 4: ALIGNMENT RESULTS			
	GB_EST1:Z14808	331	Z14808	CELSE4 Chris Martin sorted cDNA library Caenorhabditis elegans cDNA	Caenorhabditis elegans	36,890	19-Jun-97
rxa01288 498	GB_VI:S62819	3348	S62819	F2L=putative RNA polymerase-associated transcription factorF4R=type I orf virus topoisomerase homolog [orf virus OV, NZ2, host=sheep, Genomic, 3 genes,	orf virus	40,471	25-Aug-93
	GB_PR4:HUMCCLEC1 17079	1 17079	AF077344	Homo sapiens cartilage-derived C-type lectin (CLECSF1) gene, exons 1	Homo sapiens	34,631	15-OCT-1999
	GB_PR4:HUMCCLEC1 17079	1 17079	AF077344	and 2. Homo sapiens cartilage-derived C-type lectin (CLECSF1) gene, exons 1 and 2	Homo sapiens	39,300	15-OCT-1999
rxa01354 1059		301692	D87675	and 2. Homo sapiens DNA for amyloid precursor protein, complete cds.	Homo sapiens	37,984	22-Sep-97
	GB_PR1:D87675	301692		Homo sapiens DNA for amyloid precursor protein, complete cds.	Homo sapiens	35,140	22-Sep-97
	GB_RO:MMNUCLEO	11478	66920X	Mouse nucleolin gene.	Mus musculus	37,146	27-Aug-98
rxa01376 984	GB_BA1:MTCY71	42729	Z92771	Mycobacterium tuberculosis H37Rv complete genome; segment 141/162.	Mycobacterium tuberculosis	39,496	10-Feb-99
	GB_BA1:ACCPSXM	2748	X81320	A.calcoaceticus epsX and epsM genes.	Acinetobacter calcoaceticus	40,353	19-OCT-1994
	GB_BA2:ECU05248	1781	U05248	Escherichia coli polysialic acid gene cluster region 2 (neuD and neuB)	Escherichia coli	34,995	1-Feb-95
rxa01385 2004	4 GB_BA1:FVBPENTA	2519	M98557	genes, complete cds. Flavobacterium sp. pentachlorophenol 4-monooxygenase gene, complete mbNA	Flavobacterium sp.	40,855	26-Apr-93
	GR PAT-119994	2516	119994	Sections 2 from patent US 4512478	Inknown	40 855	07-OCT-1996
	GB_177.45059	2,45	AEDSOBB	Schingen E. F. Company of Control	Sphingomonas en 11630	42 993	27-Anr-99
		01+7	0000000	opfilingdrindras sp. Octor perfaction optierior 4-monooxygenase (pupp) gene, complete cds; and pentachlorophenol 4-monooxygenase reductase (pcpD) gene, partial cds.	opiningonionas sp. 0000	26,24	66-144-17
rxa01426 750	GB_GSS3:B35912	313	B35912	HS-1031-A2-D02-MR abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 811 Col=4 Row=G, genomic survey sequence.	Homo sapiens	38,019	17-OCT-1997
	GB_GSS1:FR0027767	7 497	AL020589	F.rubripes GSS sequence, clone 197B17aA3, genomic survey sequence.	Fugu rubripes	35,814	10-DEC-1997
	GB_GSS5:AQ774340	449	AQ774340		Homo sapiens	40,535	29-Jul-99
rxa01427 1044	4 GB_BA2:AF036766	3487	AF036766	sequence. Lactobacillus reuteri plasmid pTE15 replication-associated protein A (repA)	Lactobacillus reuteri	39,101	19-Feb-98
				and replication-associated protein B (repB) genes, complete cds.			
	GB_PR4:AC007032	126803		Homo sapiens clone NH0022N19, complete sequence.	Homo sapiens	34,180	17-Jul-99
	GB_PR4:AC007032	126803		Homo sapiens clone NH0022N19, complete sequence.	Homo sapiens	36,858	17-Jul-99
rxa01428 1260		41625	AL049826	Streptomyces coelicolor cosmid H24.	Streptomyces coelicolor	51,278	11-MAY-1999
	GB_BA2:AF031590	9299	AF031590	Streptomyces coelicolor thioredoxin (trxA) gene, partial cds; SpoOJ-like, Soj-Streptomyces coelicolor like, GidB-like, Jac-like inner membrane protein, and 9-10kDa protein-like.	-Streptomyces coelicolor	39,389	20-Feb-98
				genes, complete cds; RNase P protein (rnpA) gene, partial cds; and			
	MOAYOTOO. LAG GO	2733 0	716311	unknown gene.	Otrophomycon poolingler	20.280	10 057 1009
	מיסט באייאטריטטייבעם פס	0 0 L	200	Sueptoniyos coelicatol tixa a tipa genes a One's 203, 344, 233, 233, 233, 170, 341 & 124.	onepromiyes coencolor	600'60	9661-017-01
xa01430 1311	1 GB_EST30:AI643302	254	A1643302	vi39b08.y1 Stratagene mouse skin (#937313) Mus musculus cDNA clone IMAGE:974583 5' similar to SW:6PGD_HUMAN P52209 6-PHOSPHOGLUCONATE DEHYDROGENASE, DECARBOXYLATING;	Mus musculus	38,627	29-Apr-99
				HININA sequence.			

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		GB_HTG3:AC010363	174962	AC010363			35,784	15-Sep-99
rxa01465 1284	284	GB_BA1:ROX99622 GB_BA1:D83237	7224 1626	X99622 D83237	IN PROGRESS ***, 43 unordered pieces. Rhodococcus opacus catR, catA, catB, catC genes and five ORFs. Rhodococcus erythropolis DNA for catechol 1,2-dioxgenase, complete cds.	Rhodococcus opacus Rhodococcus erythropolis	58,814 53,904	24-Sep-97 1-Sep-99
		GB_EST9:AA119571	445	AA119571	mp68d04.r1 Soares 2NbMT Mus musculus cDNA clone IMAGE:574375 5' similar to TR:G559375 G559375 RAS GTPASE-ACTIVATING PROTEIN. ;; mRNA sequence	Mus musculus	39,551	17-Feb-97
rxa01466 1083	083	GB_EST37:A1934978	425	A1934978	es_NFL_T_GBC_S1 Homo sapiens cDNA clone 3. mRNA sequence.	Homo sapiens	43,609	2-Sep-99
		GB_EST15:AA465729	289	AA465729	sapiens cDNA clone IMAGE:815002	Homo sapiens	41,115	13-Aug-97
		GB_EST24:Al219091	633	AI219091	placenta_8to9weeks_2NbHP8to9W Homo sapiens 1759280 3' similar to TR:Q9988 Q99988 TGF-BETA 7TFIN 11 : mRNA sequence	Homo sapiens	36,066	29-Nov-98
rxa01477 1671	571	GB_BA2:CGU89648	1105	U89648	beacterium glutamicum unidentified sequence involved in histidine	Corynebacterium glutamicum 49,726	49,726	30-MAR-1999
		GB_EST21:AA919685	782	AA919685	vivaluesis, parier sequence. vx11g06.r1 Soares 2NbMT Mus musculus cDNA clone IMAGE:1264186 5' similar to gb:M73696 Murine Glvr-1 mRNA, complete cds (MOUSE);, mRNA	Mus musculus	37,762	20-Apr-98
		GB_HTG2:HS1005F21 101795	101795	AL078633	sequence.  Homo sapiens chromosome 20 clone RP5-1005F21, *** SEQUENCING IN PROGRESS *** in unordered pieces.	Homo sapiens	38,371	30-Nov-99
rxa01499 394	3945	GB_PR4:AC006454	153201	AC006454	Homo sapiens clone DJ0852P06, complete sequence.	Homo sapiens	38,033	13-Aug-99
		GB_BA1:LSLYSSYNT	4724	X96558			42,840	8-Jan-97
		GB_PR4:AC006454	153201	AC006454	olete sequence.	iens	38,823	13-Aug-99
rxa01502 139	1356	GB_PAT:192046	2203	192046			39,755	01-DEC-1998
		GB_PAT:178757	2203	178757	Sequence 13 from patent US 5693781.		39,755	3-Apr-98
		GB_BA1:MTCY359	36021	Z83859	complete genome; segment 84/162.	Sis	36,613	17-Jun-98
rxa01509 597	26	GB_BA1:SCE9	37730	AL049841			60,637	19-MAY-1999
		GB_BA1:MTY15C10	33050	Z95436	omplete genome; segment 154/162.	ulosis	59,296	17-Jun-98
		GB_BA1:MLCB2548	38916	AL023093	Mycobacterium leprae cosmid B2548.	n leprae	59,764	27-Aug-99
rxa01510 1404	404	GB_GSS9:AQ129927	440	AQ129927	HS_2165_B1_D09_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2165 Col=17 Row=H, genomic survey sequence.	Homo sapiens	36,136	23-Sep-98
		GB_BA2:AF016585	41097	AF016585	Streptomyces caelestis cytochrome P-450 hydroxylase homolog (nidi) gene, Streptomyces caelestis partial cds. polyketide synthase modules 1 through 7 (nidA) genes.		37,464	07-DEC-1997
			0	1	ete cds; and N-methyltransferase homolog gene, partial cds.		000	100 00 1000
		GB_HTG4:AC010747	216500	AC010747	Homo sapiens chromosome unknown clone NH0555HU9, WURKING DRAFT SEQUENCE, in unordered pieces.	nomo sapiens	33,022	29-001-1388
rxa01511 1065	990	GB_BA1:BRLBIOBA	1647	D14084	hetase, complete cds.	Corynebacterium glutamicum 40,283	40,283	3-Feb-99
		GB_GSS3:B45213	358	B45213	HS-1060-B2-D07-MF.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 782 Col=14 Row=H, genomic survey sequence.	Homo sapiens	49,505	21-OCT-1997

## TABLE 4: ALIGNMENT RESULTS

29-OCT-1999	18-Jun-98 19-Aug-98	27-Aug-99 30-Jan-96 2-Aug-97	28-Apr-93 21-MAR-1999 21-MAR-1999		23-Nov-99	6-Apr-99 04-OCT-1999	08-DEC-1999	08-DEC-1999	27-Jan-99	55-00L-51	13-Feb-99	24-Apr-93	4-Jun-98	14-Jul-99	9-Nov-97	23-Nov-99	23-Nov-99	2-Sep-99 11-MAR-1999
33,819	40,354 60,814	39,992 39,126 34,227	38,414 36,919 38,130		35,501	40,497 39,699	34,516	36,177	40,345	34,783	37,598	39,173	n 99,636	39,231	37,431	35,345	37,381	36,140 36,725
Homo sapiens	Mycobacterium tuberculosis Mycobacterium smegmatis	Mycobacterium leprae Mycobacterium leprae Caenorhabditis elegans	Discopyge ommata Vibrio cholerae Vibrio cholerae		Nomo sapiens	Ralstonia eutropha Caenorhabditis elegans	Homo sapiens	Homo sapiens	bovine adenovirus type 3	Acetobacter xylinus	Acetobacter xylinus	Acetobacter xylinus	Corynebacterium glutamicum 99,636	Helianthus annuns	Caenorhabditis elegans	Homo sapiens	Homo sapiens	Caenorhabditis elegans Homo sapiens
Homo sapiens chromosome unknown clone NH0555H09, WORKING	DRAFT SEQUENCE, in unordered pieces.  Mycobacterium tuberculosis H37Rv complete genome; segment 153/162.  Mycobacterium smegmatis cell division protein (FtsH) gene, complete cds.	Mycobacterium leprae cosmid B2548. Mycobacterium leprae cosmid B1308. Caenorhabditis elegans cosmid F27E11.	Discopyge ommata (clone OL4) agrin mRNA, 3' end cds. Vibrio cholerae Rtx toxin gene cluster, complete cds. Vibrio cholerae Rtx toxin gene cluster, complete cds.		Human DNA sequence from PAC 52D1 on chromosome Xq21. Contains CA Homo sapiens repeats. STS	Ralstonia eutropha phasin (phaP) mRNA, complete cds.  Complete cds.  Complete cds.  Complete cds.	mknA, complete cas.  Homo sapiens clone RP11-399K21, *** SEQUENCING IN PROGRESS ***, 35 unordered pieces.	45 unordered pieces.  35 unordered pieces.	Bovine adenovirus 3 complete genome.	Acetobacter xylinus genes for endoglucanase, ceilulose syntnase subunit ABCD and beta-glucosidase, complete cds.	Acetobacter xylinus genes for endoglucanase, cellulose synthase subunit ABCD and beta-glucosidase, complete cds.	A.xylinum bcs A, B, C and D genes, complete cds's.	Corynebacterium glutamicum beta C-S lyase (aecD) and branched-chain amino acid uptake carrier (brnQ) genes, complete cds, and hypothetical protein Yhbw (vhbw) gene partial cds.	H.annuus mRNA for aquaporin.	Caenorhabditis elegans chromosome V clone Y32F6, *** SEQUENCING IN PROGRESS *** in unordered pieces.	Homo sapiens Chromosome 16 BAC clone CIT987SK-625P11, complete sequence	Homo sapiens Chromosome 16 BAC clone CIT987SK-625P11, complete sequence.	Caenorhabditis elegans cosmid R11A5, complete sequence. to02d01.x1 NCI_CGAP_Ut2 Homo sapiens cDNA clone IMAGE:2177857 3' similar to SW:NU4M_PANTR P03906 NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4; mRNA sequence.
AC010747	Z95557 AF037269	AL023093 U00012 AF016413	L01423 AF119150 AF119150		Z96811	AF079155 AF039570	AC010997	AC010997	AF030154	AB010645	AB010645	M37202	M89931	X95952	AL008875	AC004125	AC004125	Z83122 Al499508
216500	24244 2364	38916 33312 25700	4354 18605 18605		148691	686 1866	187768	187768	34446	16836	16836	; <del>D</del> 9540	2821	931	187816	194020	194020	26671 403
GB_HTG4:AC010747	GB_BA1:MTCY7H7B GB_BA2:AF037269	GB_BA1:MLCB2548 GB_BA1:U00012 GB_IN1:CELF27E11	GB_OV:DYGAGR GB_BA2:AF119150 GB_BA2:AF119150		GB_PR3:HS52D1	GB_BA2:AF079155 GB_IN2:AF039570	GB_HTG7:AC010997	GB_HTG7:AC010997	GB_VI:AF030154	GB_BA1:AB010645	GB_BA1:AB010645	GB_BA1:ABCBCSABCD3540	GB_BA2:CORCSLYS	GB PL2:HAAP	GB_HTG1:CEY32F6	GB_PR4:HUAC004125 194020	GB_PR4:HUAC004125 194020	GB_IN1:CER11A5 GB_EST28:Al499508
	2682	066	1962		3441		1584			909			1098			1173		1200
	rxa01513	rxa01593	rxa01608	гха01620	rxa01640		rxa01653		0.1	rxa01/16			rxa01728			rxa01732		rxa01810 1200

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					TABLE 4: ALIGNING INCOLLE			
		GB_EST28:Al499508	403	AI499508	to02d01.x1 NCI_CGAP_Ut2 Homo sapiens cDNA clone IMAGE:2177857 3' Homo sapiens similar to SW:NU4M_PANTR P03906 NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4;, mRNA sequence.		38,264	11-MAR-1999
rxa01828 1545	1545	GB_BA1:MLCB1770 GB_HTG2:AC008073	37821 173144	Z70722 AC008073	Mycobacterium leprae cosmid B1770.  Homo sapiens clone NH0507M03, *** SEQUENCING IN PROGRESS ***, 3 Homo sapiens unordered pieces.	ı leprae	36,411 36,310	29-Aug-97 17-Jul-99
		GB_HTG2:AC008073	173144	AC008073	Homo sapiens clone NH0507M03, *** SEQUENCING IN PROGRESS ***, 3 Homo sapiens unordered pieces.		36,310	17-Jul-99
rxa01829 1446	1446	GB_IN1:AB018544 GB_EST8:AA003136	620 450	AB018544 AA003136		apillata	34,855 42,202	6-Feb-99 19-Jul-96
		OD 1814. A DOGGES	C	0000	IMAGE:42/320 5 SIMIIST to gb.X0/315 PLACENTAL PROTEIN 15 AN);, mRNA sequence.		35 Q68	6 <u>-</u> Feh.99
rxa01868 20	2049	GB_BA1:MTV033	520 21620	ABU18544 AL021928	Hydra magnipapinata mitwa for nym-179 prepromonet, compree cos. Trypia magnipapinata Mycobacterium tuberculosis H37Rv complete genome; segment 11/162. Mycobac	nyora magnipapinata Mycobacterium tuberculosis 3	38,679	17-Jun-98
		GB_BA1:MLCL622		Z95398			38,911	24-Jun-97
rxa01934 68	681	GB_BA1:MSGB983CS GB_PR4:DJ534K4	36788 216387	L78828 AF109907	Mycobacterium leprae cosmid B963 DNA sequence. Homo sapiens \$164 gene, partial cds; PS1 and hypothetical protein genes, Homo sapiens	ı ieprae	39,189	13-Jun-96 23-DEC-1998
		GB_HTG2:AC006342	201618	AC006342			34,412	11-Jan-99
		_ GB_HTG2:AC006342	201618		unordered pieces. Homo sapiens clone DJ0054D12, *** SEQUENCING IN PROGRESS ***, 3 Homo sapiens		34,412	11-Jan-99
71 7967 17	1266	GR 1N2-AC005467	62091	AC005467	unordered pieces. Drosonhija mejanogaster chromosome 2R -region 48C1-48C2, P1 clone - Drosophi	Orosophila melanogaster 3	35.252	12-DEC-1998
	2							
		GB_BA2:AE001678 GB_IN2:AC005467	13485	AE001678 AC005467	n 94 of 103 of the complete genome. mosome 2R, region 48C1-48C2, P1 clone	Chlamydophila pneumoniae 3 Drosophila melanogaster 3	35,203 34,699	08-MAR-1999 12-DEC-1998
rxa01993 1	1166	GB_BA1:PPVANAB	2864				51,697	09-MAY-1998
		GB_HTG2:AC006799	278007	AC006799	Caenorhabditis elegans clone Y51H7, *** SEQUENCING IN PROGRESS Caenorh. ***, 7 unordered pieces.	Caenorhabditis elegans 3	38,455	23-Feb-99
		GB_HTG2:AC006799	278007	AC006799	Caenorhabditis elegans clone Y51H7, *** SEQUENCING IN PROGRESS ***, 7 unordered pieces.	Caenorhabditis elegans 3	38,455	23-Feb-99
rxa01994 1098	1098	GB_HTG4:AC009961	231522	AC009961	Homo sapiens chromosome unknown clone NH0357L02, WORKING Homo sapiens DRAFT SFOLIENCE in unordered pieces.		35,576	29-OCT-1999
		GB_HTG4:AC009961	231522	AC009961	Homo sapiens chromosome unknown clone NH0357L02, WORKING Homo sapiens DRAFT SEQUENCE, in unordered pieces.		35,576	29-OCT-1999
		GB_HTG4:AC009961	231522	AC009961	Homo sapiens chromosome unknown clone NH0357L02, WORKING Homo sapiens DRAFT SEQUENCE, in unordered pieces.		35,472	29-OCT-1999
rxa01997 60	609	GB_BA2:AF112536	1798	AF112536	te reductase beta-chain (nrdF)	Corynebacterium glutamicum 37,719	37,719	5-Aug-99
		GB_BA1:SCH66 GB_EST29:AI558691	9153 598	AL049731 Al558691	omyces coelicolor cosmid H66.  10.y1 Zebrafish WashU MPIMG EST Danio rerio cDNA 5' similar to	es coelicolor	38,655 40,232	29-Apr-99 24-MAR-1999
					SW:ATP_HOMAN P1884/ CTCLIC-AMP-DEPENDENT TRANSCRIPTION FACTOR ATF-3; mRNA sequence.			

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26-MAY-1995	42 Con 07	20-Jun-98	8-Jul-99	4-Sep-98	22-Aug-94	15-Jun-96	17-Jun-98	02-OCT-1998	09-OCT-1999	02 OCT 4007	09-OCT-1999		8-Aug-98	23-Nov-99	16-MAY-1996	14-MAY-1999	14-MAY-1999	10-Nov-97	30-Aug-99		07-MAY-1999	24-MAR-1999	01-MAR-1994	24-Jun-99	7	20-MAR-1998	23-Apr-99
35,920		35,599	32,935	32,935	31,995	50,604	38,113	41,876	36,818	97 547	37,517 35,563		36,149	37,587	36,755	99,766	36,983	37,231	37,500		38,268	50,791	37,563	39,504 38 377		39,387	38,377
Homo sapiens		Homo sapiens	Homo sapiens	Homo sapiens	Homo sapiens	Mycobacterium leprae	Mycobacterium tuberculosis	Ochrobactrum anthropi	Homo sapiens		Arabidopsis mana Homo sapiens			Homo sapiens	Homo sapiens	Corynebacterium glutamicum 99,766	Corynebacterium glutamicum 36,983	Citrullus lanatus	Homo sapiens		Schizosaccharomyces pombe	myces coelicolor		Mycobacterium tuberculosis	200485	Homo sapiens	Homo sapiens
TABLE 4: ALIGNMENT RESULTS vi18d08.r1 Soares placenta Nb2HP Homo sapiens cDNA clone	IMAGE:139599 5', mRNA sequence.	CIT-HSP-385H2. TRB CIT-HSP Homo sapiens genomic clone 385H2, complete sequence.	Homo sapiens interferon-induced protein p78 (MX1) gene, complete cds.	Homo sapiens chromosome 21, P1 clone LBL#8 (LBNL H8), complete sequence	Homo sapiens (subclone H8 10_f11 from P1 35 H5 C8) DNA sequence.	Mycobacterium leprae cosmid B32 DNA sequence.	Mycobacterium tuberculosis H37Rv complete genome; segment 127/162.	3123-4r Ochrobactrum anthropi BAC Library Ochrobactrum anthropi	genomic clone 3l23-4r, genomic survey sequence. Homo sapiens 12q22-103.4-106.5 BAC RPCI11-718L23 (Roswell Park	Cancer Institute Human BAC Library) complete sequence.	Genomic sequence of Afablopsis BAC FoA5, complete sequence. Homo sapiens 12o22-103 4-106 5 BAC RPCI11-718L23 (Roswell Park	Cancer Institute Human BAC Library) complete sequence.	Lactobacillus helveticus cochaperonin GroES and chaperonin GroEL genes, Lactobacillus helveticus complete cds; and DNA mismatch repair enzyme (hexA) gene, partial cds.	Homo sapiens chromosome 6 clone RP1-34F7, *** SEQUENCING IN PROGRESS ***, in unordered pieces.	Human RP1 and complement C4B precursor (C4B) genes, partial cds.	Corynebacterium glutamicum amtP, glnB, glnD genes and partial ftsY and srp genes.	Corynebacterium glutamicum amtP, glnB, glnD genes and partial ftsY and srp genes.	EST00115 watermelon lambda zap express library Citrulius lanatus cDNA	cione www.5233 5 similar to transiation initiation factor, mixiva sequence. HS_4640_B2_F09_T7A CIT Approved Human Genomic Sperm Library D	Homo sapiens genomic clone Plate=4640 Col=18 Row=L, genomic survey sequence.	S.pombe chromosome III cosmid c970.	Streptomyces coelicolor cosmid 6G10.	Mycobacterium leprae cosmid B1170.	Mycobacterium tuberculosis H37Rv complete genome; segment 70/162.  Homo canians clone NH0304410 *** SEOLIENCING IN PROGRESS *** 3		Homo sapiens PAC clone DJ0789N01 from 7q21, complete sequence.	Homo sapiens clone NH0304A10, *** SEQUENCING IN PROGRESS ***, 3 unordered pieces.
R64206	00000	AC002340 B55001	AF135187	AC005612	L35666	L78818	Z74697	AQ242118	AC008055	00000	AC008055		AF031929	AL049547	U24578	AJ010319	AJ010319	AA660065	AQ839377		AL031530	AL049497	U00010	Z95586 AC007164		AC004451	AC007164
453	1006	406	33016	60904	3949	36404	29372	. 992	196899	120707	196899		2675	129811	17488	5368	5368	352	523		31438	36734	41171	32437	0000	108642	158320
GB EST3:R64206		GB_GSS3:B55001	GB PR4:AF135187	GB_PR3:AC005612	GB_PR1:HUM8DC11Z	GB_BA1:MSGB32CS	GB_BA1:MTCY338	GB_GSS10:AQ242118	GB_PR4:AC008055	200000	GB_PR4:AC002232 GB_PR4:AC008055		GB_BA2:AF031929	GB_HTG1:HSDJ34F7	GB_PR2:HSU24578	GB_BA1:CAJ10319	GB_BA1:CAJ10319	GB_EST17:AA660065	GB_GSS6:AQ839377		GB_PL1:SPCC970	GB_BA1:SC6G10	GB_BA1:U00010	GB_BA1:MTCY336 GB_HTG2:AC007164		GB_PR3:AC004451	GB_H1GZ:AC0071b4
915			1 762			3010			3 1533				1761		:	1869		391				1407		465			
rxa02052			rxa02064			rxa02082			xa02083				rxa02092			rxa02098		rxa02105				rxa02111		0xa02118	700		

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					TABLE 4: ALIGNMENT RESULTS			
xa02448 1212	1212	GB_BA1:AB016258	, 0922	AB016258	e reductase and hydroxyquinol 1,2-	Arthrobacter sp. 6	60,465	8-Sep-99
		GB_EST37:AW014148 !	553	AW014148	oloxygenase, partial and complete cus. UI-H-BIQ-aaj-c-04-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone เหลดกะววกดุผลว 3' mDNA รอกแอกกล	Homo sapiens 4	44,560	10-Sep-99
		GB_EST14:AA432042	543	AA432042	sapiens cDNA clone IMAGE:782521 mRNA sequence	Homo sapiens 3	36,522	22-MAY-1997
rxa02449 1026	1026	GB_BA1:AB016258	, 2260	AB016258	s reductase and hydroxyquinol 1,2-	Arthrobacter sp. 6	66,244	8-Sep-99
		GB_BA1:CGPUTP	3791	Y09163	dioxygenase, panial and complete cas. C.glutamicum putP gene.	Corynebacterium glutamicum 39,899	39,899	8-Sep-97
		GB_BA1:AB016258	, 2260	AB016258	Arthrobacter sp. gene for maleylacetate reductase and hydroxyquinol 1,2-	Arthrobacter sp. 7	70,410	8-Sep-99
rxa02497 1	1050	GB_BA2:CGU31224	422	U31224	ie, partial cds.	Corynebacterium glutamicum 96,445	96,445	2-Aug-96
		20G9	37218	277162	complete genome; segment 25/162.	osis	59,429	17-Jun-98
rxa02526 1329	1329	က	16911 483	AL049819 AQ240233	Streptomyces coelicolor cosmid E7. CIT-HSP-2385F9.TR.1 CIT-HSP Homo sapiens genomic clone 2385F9,	Streptomyces coelicolor 3 Homo sapiens 4	39,510 42,475	10-MAY-1999 30-Sep-98
			195	S48556	genomic survey sequence. {tandem repeat P1 monomer} {Cacatua galerita=sulfur-crested cockatoo,	Cacatua galerita	50,515	08-MAY-1993
		GB_PR2:HSM801056	2555	AL117532	Genomic, 133 mg. Homo sapiens mRNA; cDNA DKFZp434E192 (from clone DKFZp434E192). Homo sapiens		39,116	15-Sep-99
rxa02530 780	780	GB_PR3:HSJ753D10 (	97912	AL049651	Human DNA sequence from clone 753D10 on chromosome 20 Contains genes for SSTR4(somatostatin receptor 4) and THBD(thrombomodulin),	Homo sapiens	34,248	23-Nov-99
		GB_EST33:AI782764 (	661	AI782764	ulentum cDNA	Lycopersicon esculentum 3	35,385	29-Jun-99
		GB_GSS9:AQ121479	521	AQ121479	HS_3084_A2_B02_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3084 Col=4 Row=C, genomic survey	Homo sapiens	38,689	22-Sep-98
rxa02535 1278	1278	GB_HTG3:AC008710	146065	AC008710	sequence.  Homo sapiens chromosome 5 clone CIT978SKB_7E3, *** SEQUENCING IN DEOCRESS *** 30 unordered pieces	Homo sapiens	35,799	3-Aug-99
		GB_HTG3:AC008710	146065	AC008710	Homo sapiens chromosome 5 clone CIT978SKB_7E3, *** SEQUENCING IN PROCEES, *** SEQUENCING	Homo sapiens	35,799	3-Aug-99
		GB_HTG3:AC008710	146065	AC008710	78SKB_7E3, *** SEQUENCING	Homo sapiens	34,886	3-Aug-99
rxa02603 1119	1119	GB_BA1:MTV026 GB_IN2:AC005714	23740 177740	AL022076 AC005714	Mycobacterium tuberculosis H37Rv complete genome; segment 157/162. Drosophila melanogaster, chromosome 2R, region 58D4-58E2, BAC clone	Mycobacterium tuberculosis 3 Drosophila melanogaster 4	37,975 41,226	24-Jun-99 01-MAY-1999
		GB_EST19:AA775050	218	AA775050	DACK40M 13, Complete sequence. ac76e10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:868554 3' similar to gb:Y00371_rna1 HEAT SHOCK COGNATE 71 KD PROTFIN (HIMAN): mRNA sequence.	Homo sapiens	40,826	5-Feb-98
rxa02641								

	17-Jun-98 5-Aug-99	27-Aug-99 13-MAY-1999	19-MAY-1999 13-MAY-1999	6-Feb-99	4-Aug-97	08-OCT-1998	6-Feb-99			07-OCT-1996 08-OCT-1997	(Rel. 52, Created)			08-0C1-199/ (Rel. 52, Created)	10-Feb-99 08-OCT-1997 (Rel 52	
	39,109	62,753 58,095	38,544	m 99,365	m 99,317	m 99,296	m 97,468			97,619 m 97,619		37,856	98,605	c09,88 ⊞	s 34,868 im 98,547	98,547
	Mycobacterium tuberculosis Streptomyces coelicolor A3(2)	Mycobacterium leprae Pseudomonas putida	Streptomyces coelicolor Pseudomonas putida	Corynebacterium glutamicum 99,365	Corynebacterium glutamicum 99,317	Corynebacterium glutamicum 99,296	Corynebacterium glutamicum 97,468	Vicia faba Bacillus sp. Mycobacterium tuberculosis	Archaeoglobus fulgidus Mycobacterium tuberculosis	Unknown. 97,619 Corynebacterium glutamicum 97,619		Streptomyces coelicolor A3(2)	Unknown.	Corynebacterium glutamicum 98,605	Mycobacterium tuberculosis 34,868 Corynebacterium glutamicum 98,547	Unknown.
TABLE 4: ALIGNMENT RESULTS	Mycobacterium tuberculosis H37Rv complete genome; segment 69/162. Streptomyces coelicolor cosmid 4A10.	Mycobacterium leprae cosmid L458.  Pseudomonas putida NCIMB 9866 plasmid pRA4000 p-cresol degradative pathway genes, p-hydroxybenzaldehyde dehydrogenase (pchA), p-cresol methylhydroxylase, cytochrome subunit precursor (pchC), unknown (pchX) and p-cresol methylhydroxylase, flavoprotein subunit (pchF) genes, complete cds.	Streptomyces coelicolor cosmid E9. Pseudomonas putida NCIMB 9869 plasmid pRA500 p-cresol degradative pathway genes, p-hydroxybenzaldehyde dehydrogenase (pchA) gene, partial cds, and p-cresol methylhydroxylase, cytochrome subunit (pchC), unknown (pchX), p-cresol methylhydroxylase, flavoprotein subunit (pchF), protocatechuate-3,4-dioxygenase, beta subunit (pcaH) and protocatechuate-3,4-dioxygenase, alpha subunit (pcaG) genes, complete cds.	Corynebacterium glutamicum genes for MurC and FtsQ, complete cds.	Corynebacterium glutamicum gene for MurC, FtsQ, FtsZ, complete cds.	B.lactofermentum murC, ftsQ or divD & ftsZ genes.	Corynebacterium glutamicum genes for MurC and FtsQ, complete cds.	V.faba mRNA for amino acid transporter. DNA encoding recombinant monoglyceride lipase. Mycobacterium tuberculosis H37Rv complete genome; segment 96/162.	Archaeoglobus fulgidus section 146 of 172 of the complete genome. Mycobacterium tuberculosis H37Rv complete genome; segment 96/162.	Sequence 4 from patent US 5556776. Base sequence of sucrase gene.		Streptomyces coelicolor cosmid 4A10.	Sequence 4 from patent US 5556776.	Base sequence of sucrase gene.	Mycobacterium tuberculosis H37Rv complete genome; segment 96/162. Base sequence of sucrase gene.	Sequence 4 from patent US 5556776.
	Z74020 AL109663	AL049478 U96338	AL049841 U96339	AB015023	AB003132	Y08964	AB015023	Y09591 E05047 Z95388	AE000961 Z95388	126124 E11760		AL109663	126124	E11760	Z95388 E11760	126124
	35377 43147	43839 5276	37730 4464	2291	4116	5546	2291	. 1879 966 37586	18765 37586	6911		43147	6911	6911	37586 6911	6911
	GB_BA1:MTCY48 GB_BA1:SC4A10	GB_BA1:MLCL458 GB_BA2:PPU96338	GB_BA2:PPU96339 GB_BA2:PPU96339	GB_BA1:AB015023	GB_BA1:AB003132	GB_BA1:BLFTSZ	GB_BA1:AB015023	GB_PL2:VFAMACTRA 1 GB_PAT:E05047 GB_BA1:MTCY270	GB_BA2:AE000961 GB_BA1:MTCY270	GB_PAT:126124 EM_PAT:E11760	l	GB_BA1:SC4A10	GB_PAT:126124	EM_PAT:E11760	GB_BA1:MTCY270 EM_PAT:E11760	GB_PAT:126124
	1053	1575		1581			1212.	1812	!	1539			1221		1653	
	rxa02651 1053	ка02674 1575		rxa02702			rxa02703 1212.	rxa02704		rxa02705			rxa02706 1221		rxa02707 1653	

Table 4, Page 16

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TABLE 4: ALIGNMENT RESULTS

rxa02710 1686	GB_BA1:MLCB268 EM_PAT:E11760	38859 6911	AL022602 E11760	Mycobacterium leprae cosmid B268. Base sequence of sucrase gene.	Mycobacterium leprae 37,815 Corynebacterium glutamicum 52,124	37,815 52,124	27-Aug-99 08-OCT-1997 (Rel. 52,
	GB_PAT:126124 GB_GSS13:AQ484169	6911 9 515	126124 AQ484169	Sequence 4 from patent US 5556776. RPCI-11-264A12.TV RPCI-11 Homo sapiens genomic clone RPCI-11-264A12 genomic clone RPCI-11-	Unknown. 52 Homo sapiens · 40	52,124 40,856	Oreated) 07-OCT-1996 24-Apr-99
rxa02711 2235	GB_BA2:XCU45994	1203	U45994	Xanthomonas campestris pv. campestris insertion sequence IS1404.	Xanthomonas campestris pv. 36	39,061	29-Jan-99
	GB_BA2:XCU77781	4160	U77781	Xanthomonas campestris pv. amaranthicola Xaml DNA methyltransferase (xamlM) gene, complete cds; insertion sequence IS1389 and unknown	Camposins Xanthomonas campestris pv. 39,551 amaranthicola	39,551	9-Feb-99
	GB_BA2:AF108355	1222	AF108355	Yerres.  Xanthomonas campestris pv. amaranthicola insertion sequence IS1389-B	Xanthomonas campestris pv. 40,281	10,281	09-MAR-1999
rxa02713 1134	GB_BA1:MTCY270	37586	Z95388	Mycobacterium tuberculosis H37Rv complete genome; segment 96/162.	n tuberculosis	38,669	10-Feb-99
	GB_PR1:D31907 GB_PR1:HSMTFMR	599 3302	D3190/ X78710	Homo sapiens gene for zinc regulatory factor, partial cds. H.sapiens MTF-1 mRNA for metal-regulatory transcription factor.	Homo sapiens 37	30,390 37,243	7-reb-99 1-Aug-94
rxa02716 684	GB_PR3:AC002347 GB_PR3:HS310J6	134977 87942	AC002347 AL035593	Homo sapiens chromosome 17, clone 297N7, complete sequence.  Human DNA sequence from clone 310J6 on chromosome 6q22.1-22.3.	Homo sapiens 30 Homo sapiens 3	36,282 37,291	3-Feb-98 23-Nov-99
				Contains part of a novel gene, ESTS, STSS and GSSS, complete sequence.			
	GB_HTG3:AC011509		111353 AC011509	Homo sapiens chromosome 19 clone CITB-H1_2189E23, *** ceru ienicinic in Deocesce *** 34 mondared pieces	Homo sapiens 3.	37,407	07-OCT-1999
rxa02722 1449	GB_BA1:BLFTSZ	5546	Y08964	SEQUENCING IN TROCKESS , 33 disordered preces.  B.lactofermentum murC, ftsQ or divD & ftsZ genes.	Corynebacterium glutamicum 99,652	99,652	08-OCT-1998
	GB_BA1:AB003132	4116	AB003132	Corynebacterium glutamicum gene for MurC, FtsQ, FtsZ, complete cds.	Corynebacterium glutamicum 98,535	38,535	4-Aug-97
	GB_PAT:E17182	1125	E17182	Brevibacterium flavum ftsQ gene complete cds.	Corynebacterium glutamicum 97,235	97,235	28-Jul-99
rxa02723 789	GB_BA1:AB015023	2291	AB015023	Corynebacterium glutamicum genes for MurC and FtsQ, complete cds.	Corynebacterium glutamicum 99,113	99,113	6-Feb-99
	GB_BA1:BLFTSZ	5546	Y08964	B.lactofermentum murC, ftsQ or divD & ftsZ genes.	Corynebacterium glutamicum 99,113	99,113	08-OCT-1998
	GB_BA1:AB003132	4116	AB003132	Corynebacterium glutamicum gene for MurC, FtsQ, FtsZ, complete cds.	Corynebacterium glutamicum 99,113	99,113	4-Aug-97
rxa02813 1108	GB_HTG3:AC009658	171795	AC009658	Homo sapiens chromosome 15 clone 344_A_16 map 15, *** SEQUENCING Homo sapiens		34,622	01-OCT-1999
	GB_HTG3:AC009658	171795	AC009658	IN PROGRESS ***, 29 unordered pieces.  Homo sapiens chromosome 15 clone 344_A_16 map 15, *** SEQUENCING Homo sapiens.  IN PROCRESS *** 30 unordered signs?		34,622	01-OCT-1999
rxa02820 1411	GB_RO:MMU65079 GB_BA1:BFU64514	2300 3837	U65079 U64514	IN PROCKESS (28 utionizated places).  Mus musculus actin-binding protein (ENC-1) mRNA, complete cds.  Bacillus firmus dppABC operon, dipeptide transporter protein dppA gene, partial cds, and dipeptide transporter proteins dppB and dppC genes.	Mus musculus Bacillus firmus	35,013 36,859	29-Jul-97 1-Feb-97
	GB_IN1:CET04C10	20958	Z69885	complete cds. Caenorhabditis elegans cosmid T04C10, complete sequence.	Caenorhabditis elegans 3	35,934	2-Sep-99

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				TIMESON AND THE THEORY AND THE THEORY			
	GB_EST35:AI823090 720	720	AI823090	L30-944T3 Ice plant Lambda Uni-Zap XR expression library, 30 hours NaCl Mesembryanthemum treatment Mesembryanthemum crystallinum cDNA clone L30-944 5' similar crystallinum to 60S ribosomal protein L36 (AC004684)[Arabidopsis thaliana], mRNA sequence.		35,770	21-Jul-99
rxa02828 572	GB_BA1:MTCY10H4 39160 Z80233 GB_BA1:MTORIREP 8400 X92504	39160 8400	Z80233 X92504	Mycobacterium tuberculosis H37Rv complete genome; segment 2/162. M.tuberculosis origin of replication and genes rnpA, rpmH, dnaA, dnaN,	Mycobacterium tuberculosis 39,823 Mycobacterium tuberculosis 39,823	39,823 39,823	17-Jun-98 26-Aug-97
rxa02839 470	GB_RO:RATENDOGLY 3906 L37380 GB_BA2:ECOUW89 176195 U00006 GB_BA2:AE000477 11314 AE000477	.Y 3906 176195 11314	3906 L37380 176195 U00006 11314 AE000477	recr. Rat apical endosomal glycoprotein mRNA, complete cds. E. coli chromosomal region from 89.2 to 92.8 minutes. Escherichia coli K-12 MG1655 section 367 of 400 of the complete genome.	Rattus norvegicus Escherichia coli Escherichia coli	38,704 99,362 99,787	20-Apr-95 17-DEC-1993 12-Nov-98
rs03218	GB_BA1:ECOPLSB	3865	K00127	E.coli pIsB and dgk genes coding for sn-glycerol-3-phosphate acyltransferase and diglyceride kinase.	Escherichia coli	33,761	28-Feb-94